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PA(IT COOPERATION TREAT(

From the INTERNATIONAL BUREAU

PCT NOTIFICATION OF ELECTION (PCT Rule 61.2)

Commissioner
US Department of Commerce
United States Patent and Trademark
Office, PCT
2011 South Clark Place Room
CP2/5C24
Arlington, VA 22202

Date of mailing: 01 November 2001 (01.11.01)	ETATS-UNIS D'AMERIQUE in its capacity as elected Office
International application No.: PCT/GB00/01675	Applicant's or agent's file reference: P23847A/JMK
International filing date: 02 May 2000 (02.05.00)	Priority date: 01 May 1999 (01.05.99)
Applicant: ADDISON, Paul, Stanley et al	

1.	The designated Office is hereby notified of its election made:
	X in the demand filed with the International preliminary Examining Authority on:
	30 November 2000 (30.11.00)
	in a notice effecting later election filed with the International Bureau on:
2.	The election X was
	was not
	made before the expiration of 19 months from the priority date or, where Rule 32 applies, within the time limit under Rule 32.2(b).
	·

The International Bureau of WIPO 34, chemin des Colombettes 1211 Geneva 20, Switzerland

Authorized officer:

J. Zahra

Facsimile No.: (41-22) 740.14.35

Telephone No.: (41-22) 338.83.38

PACINT COOPERATION TREAT

From the INTERNATIONAL BUREAU

PCT

COMMUNICATION OF INTERNATIONAL APPLICATIONS

(PCT Article 20)

Date of mailing:

26 September 2001 (26.09.01)

To

Commissioner
US Department of Commerce
United States Patent and Trademark
Office, PCT
2011 South Clark Place Room
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Arlington, VA 22202
ETATS-UNIS D'AMERIQUE

in its capacity as designated Office

The International Bureau transmits herewith copies of the international applications having the following international application numbers and international publication numbers:

International application no.:

International publication no.:

PCT/GB00/01675

The International Bureau of WIPO 34, chemin des Colombettes 1211 Geneva 20, Switzerland

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COMMUNICATION IN CASES FOR WHICH NO OTHER FORM IS APPLICABLE

MURGITROYD & COMPANY 373 Scotland Street Glasgow G5 8QA ROYAUME-UNI

Date of mailing (day/month/year) 26 September 2001 (26.09.01)					
Applicant's or agent's file reference P23847A/JMK	REPLY DUE see paragraph 1 below				
International application No. PCT/GB00/01675	International filing date (day/month/year) 02 May 2000 (02.05.00)				
Applicant THE COURT OF NA	APIER UNIVERSITY				
1. REPLY DUE within months/days from the NO REPLY DUE, however, see below IMPORTANT COMMUNICATION INFORMATION ONLY	above date of mailing				
2. COMMUNICATION:					
Due to a clerical error, the international application has not been published promptly after the expiration of 18 months from the priority date, as provided in Article 21(2) of the PCT. Consequently, the international publication will only take place on 01 November 2001 (01.11.01). Meanwhile, the International Bureau (WO) will communicate the international application to each designated Office, in accordance with Article 20. A copy of this notification has been sent to the receiving Office (RO/GB), the International Searching Authority (ISA/EP) and all the designated Offices concerned.					
The International Bureau of WIPO 34, chemin des Colombettes	Authorized officer Idhir BRITEL				

Telephone No. (41-22) 338.83.38

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Facsimile No. (41-22) 740.14.35



2	AUG 2001	
WIPO	PCT	

INTERNATIONAL PRELIMINARY EXAMINATION REPORT

(PCT Article 36 and Rule 70)

Applicant's	or age	ent's file reference		See Notification of Transmittal of International
P23847A	VER/	VPPP	FOR FURTHER ACTION	Preliminary Examination Report (Form PCT/IPEA/416)
Internationa	al appl	ication No.	International filing date (day/month	th/year) Priority date (day/month/year)
PCT/GB0	<u> </u>	675	02/05/2000	01/05/1999
G06F17/		nt Classification (IPC) or n	national classification and IPC	
Applicant THE COI	URT	OF NAPIER UNIVER	ISITY et al.	
and is	s trans	smitted to the applicant	according to Article 36.	ed by this International Preliminary Examining Authority
2. This F	REPU	HI CONSISTS OF A LOTAL O	of 12 sheets, including this cover	Sfieet.
b (s	een a see R	mended and are the ba	asis for this report and/or sheets on the Administrative Instruction	he description, claims and/or drawings which have containing rectifications made before this Authority cions under the PCT).
3. This r	_		lating to the following items:	
		Basis of the report		
!!	L] ⊠	•		westive stan and industrial applicability
III IV				ventive step and industrial applicability
v	Ø	Reasoned statement u		novelty, inventive step or industrial applicability;
VI		Certain documents ci	ted	
VII	\boxtimes	Certain defects in the	international application	
VIII	×	Certain observations of	on the international application	
Date of sub	missic	on of the demand	Date of	f completion of this report
30/11/20	00		17.08.20	2001
	exami Euro	g address of the internation ining authority: opean Patent Office	al Authoriz	ized officer
<i>())</i>		0298 Munich +49 89 2399 - 0 Tx: 52365	Barba,	ı, M
		: +49 89 2399 - 4465	Tolopho	200 No. : 40 90 2200 2722



International application No. PCT/GB00/01675

I. Basis of the report

1.	With regard to the elements of the international application (Replacement sheets which have been furnished to the receiving Office in response to an invitation under Article 14 are referred to in this report as "originally filed" and are not annexed to this report since they do not contain amendments (Rules 70.16 and 70.17)): Description, pages:								
	1-4	0	as originally filed						
	Cla	ims, No.:							
	1-4	0	as originally filed						
	Dra	wings, sheets:							
	1/14	4-14/14	as originally filed						
2.	Witl lanç	h regard to the lang guage in which the	guage, all the elements marked above were available or furnished to this Authority in the international application was filed, unless otherwise indicated under this item.						
	The	These elements were available or furnished to this Authority in the following language: , which is:							
		the language of a	translation furnished for the purposes of the international search (under Rule 23.1(b)).						
		the language of pu	ublication of the international application (under Rule 48.3(b)).						
		the language of a 55.2 and/or 55.3).	translation furnished for the purposes of international preliminary examination (under Rule						
3.			eleotide and/or amino acid sequence disclosed in the international application, the yexamination was carried out on the basis of the sequence listing:						
		contained in the in	ternational application in written form.						
		filed together with	the international application in computer readable form.						
		furnished subsequ	ently to this Authority in written form.						
		furnished subsequ	ently to this Authority in computer readable form.						
			t the subsequently furnished written sequence listing does not go beyond the disclosure in pplication as filed has been furnished.						
		The statement tha listing has been fu	t the information recorded in computer readable form is identical to the written sequence rnished.						
4.	The	amendments have	e resulted in the cancellation of:						
		the description,	pages:						
	П	the claims	Nos:						



International application No. PCT/GB00/01675

		the drawings,	sheets:										
5.		This report has been considered to go bey						nts had	not been	made,	since t	they have	been
		(Any replacement sh report.)	eet containing	such	amendn	nents n	nust be	referre	d to unde	er item :	1 and a	nnexed t	o this
6.	Add	itional observations, it	necessary:		-		-						
III.	Nor	n-establishment of op	oinion with re	gard 1	o novel	lty, inv	entive :	step ar	nd indus	trial ap	plicabi	ility	
1.		questions whether the ious), or to be industri the entire internationa	ally applicable							entive st	tep (to	be non-	
	×	claims Nos. 16-32, 3	3-40.										
be	caus	e:			•								
		the said international not require an interna						to the	following	subject	t matte	r which de	oes
	⊠	the description, claim unclear that no mean see separate sheet						s below) or said	claims l	Nos. 16	6, 38 are	so
		the claims, or said cla	ims Nos. are	so ina	adequate	ely sup	ported I	by the o	descriptic	n that r	no mea	ningful op	oinion
	×	no international searc	h report has I	oeen e	stablish	ed for t	he said	l claims	Nos. 17	-32, 39-	40.		
2.	and	eaningful internationa /or amino acid sequer ructions:											
		the written form has r	ot been furnis	shed o	r does n	ot com	ply with	the sta	andard.				
		the computer readab	e form has no	t beer	furnish	ed or d	oes not	t compl	y with the	standa	ard.		
	cita	soned statement un tions and explanatio			_		ovelty,	inventi	ive step	or indu	strial a	applicabi	lity;
1.		ement											
	Nov	elty (N)	Yes: Cl	aims	3-15, 38	5-37							



International application No. PCT/GB00/01675

No: Claims 1, 2, 33-34

Inventive step (IS) Yes: Claims

No: Claims 3-15, 35-37

Industrial applicability (IA) Yes: Claims 1-15, 33-37

No: - Claims

2. Citations and explanations see separate sheet

VII. Certain defects in the international application

The following defects in the form or contents of the international application have been noted: see separate sheet

VIII. Certain observations on the international application

The following observations on the clarity of the claims, description, and drawings or on the question whether the claims are fully supported by the description, are made: see separate sheet

Reference is made to the following documents:

D1: WO 96 08992 A (SHOSHAN HERBERT Z ;UNIV RAMOT (IL); AKSELROD SOLANGE (IL); KESELBR) 28 March 1996 (1996-03-28)

D3: SAVA H ET AL: 'APPLICATION OF THE MATCHING PURSUIT METHOD FOR STRUCTURAL DECOMPOSITION AND AVERAGING OF PHONOCARDIOGRAPHIC SIGNALS' MEDICAL AND BIOLOGICAL ENGINEERING AND COMPUTING, GB, PETER PEREGRINUS LTD. STEVENAGE, vol. 36, no. 3, 1 May 1998 (1998-05-01), pages 302-308, XP000751653 ISSN: 0140-0118

The following document (D) was not cited in the international search report. A copy of the document is annexed to the Written Opinion and the numbering will be adhered to in the rest of the procedure:

D2: Proceedings of Computers in Cardiology, IEEE, September 23-26 1991, Venice Italy, pages 393-396, D. Morlet et al: "Time-Scale Analysis of High Resolution Signal Averaged Surface ECG Using Wavelet Transformation"

- With regard to present claims 17 to 32 and 39 to 40, it is noted that no 0 international search report has been established in respect of the above mentioned set of claims. Consequently this International Preliminary Examining Authorithy, under the provisions of Rule 66.1 (e), does not need to carry out an international preliminary examination in respect to the subject matter of present claims 17 to 32 and 39 to 40.
- This International Preliminary Examination will be therefore limited to the subject matter of present claims 1 to 16 and 33 to 38.

Re Item III

Non-establishment of opinion with regard to novelty, inventive step and industrial applicability

When considering the extent of the clarity problems regarding present independent claims 16 and 38 (see Item VIII below) this International Preliminary Examining Authority considers as not possible to give an opinion as to novelty, inventive step and industrial applicability in respect of the above mentioned claim 16 and 38.

Re Item V

Reasoned statement under Rule 66.2(a)(ii) with regard to novelty, inventive step or industrial applicability; citations and explanations supporting such statement

- Insofar as the present text of independent claim 1 can be understood in the light of the description (see Item VIII below), it appears that the subject matter of claim 1 does not fulfill the requirements of novelty as set out in Article 33 (2) PCT, the reasons therefor being the following.
- 2.1 Document D1, that provisionally is considered as representing the closest prior art, discloses (see from page 9 line 27 to page 12 line 12; from page 13 line 29 to page 15 line 12) a method to analyse an ECG signal by using wavelet decomposition of said ECG signal.
 - Thus, the subject matter of claim 1 is not novel against the method known from D1 (Article 33 (2) PCT).
- Dependent claim 2 does not contain any features which, in combination with the features of any claim to which it refers, meet the requirements of the PCT in respect of novelty, because the feature of using discrete wavelet transforms is also disclosed in document D1 (Article 33 (2) PCT).

INTERNATIONAL PRELIMINARY **EXAMINATION REPORT - SEPARATE SHEET**

- Insofar as the present text of dependent claims 3 to 15 can be understood in the 4 light of the description (see Item VIII below), it also appears that the subject matter of claims 3 to 15 does not involve an inventive step and therefore the requirements of Article 33 (3) PCT are not met, the reasons therefor being the following.
- Document D2 discloses (see page 393 left column lines 1 to 21; from page 393 4.1 right column line 16 to page 394 left column line 1; page 394 right column lines 1 to 23; figure 4 and figure 5) a method to analyse an ECG signal using continuous wavelet decomposition including the following features:
 - computing wavelet energy surfaces of said ECG signal and plotting said wavelet energy surfaces against parameters a and b of the wavelets bases;
 - constructing a contour plot and a surface plot to display said wavelet ii) decomposition of said ECG signal;
 - constructing a 2D or 3D energy scalogram to display said wavelet iii) decomposition of said ECG signal.
- 4.1a Moreover, the subject matter of dependent claims 7, 8, 9, 10, 11 and 15 are considered obvious.
- 5 Therefore, the subject matter of dependent claims 3 to 15 does not include an inventive step contribution in respect of the method known from the combination of the method of D1 and the method of D2 (Article 33 (3) PCT).
- The below mentioned lack of clarity notwithstanding (see Item VIII below), the 6 subject-matter of claim 33 is not novel in the sense of Article 33 (2) PCT because document D3 discloses (see from page 303 left column line 11 to page 305 right column line 54) a method to decompose a cardiac signal using a matching pursuit algorithm.
- 7 The below mentioned lack of clarity notwithstanding (see Item VIII below), the subject-matter of independent apparatus claim 34 is not novel in the sense of Article 33 (2) PCT for the same reasons, mutatis mutandis already mentioned in

paragraph 2.1 of this International Preliminary Examination Report.

- The below mentioned lack of clarity notwithstanding (see Item VIII below), the 8 subject-matter of dependent claims 35 to 37 is not inventive in the sense of Article 33 (3) PCT for the same reasons, mutatis mutandis, already mentioned in paragraph 3 to 4.1b of this International Preliminary Examination Report.
- With regard to the assessment of the present claims 1 to 15 and 33 to 37 on the 9 question whether they are industrially applicable, the following is stated. The below mentioned lack of clarity notwithstanding (see Item VIII below), it appears that the subject matter of present claims 1 to 16 and 33 to 37 relates to a method and apparatus to decompose waveforms of a cardiac signal, therefore it fulfills the requirements of industrial applicability as set out in Article 33 (4) PCT.

Re Item VII

Certain defects in the international application

- 10 At page 6 line 28 of the description the wording "me" should be amended as "method".
 - At page 9 line 17 of the description the wording "scrologram" should be amended as "scalogram".
- 10.1 Present independent claims are not in the two-part form in accordance with Rule 6.3(b) PCT, which in the present case would be appropriate, with those features known in combination from the prior art (document D1) being placed in a preamble (Rule 6.3(b)(i) PCT) and with the remaining features being included in a characterising part (Rule 6.3(b)(ii) PCT).

Independent claims should therefore be redrafted accordingly.

In addition, it should be clear from the description which features of the claimed subject-matter are known from documents D1, D2 and D3 (see the PCT Guidelines PCT/GL/3 III, 2.3a).



- 10.2 Contrary to the requirements of Rule 5.1(a)(ii) PCT, the relevant background art disclosed in the documents D1, D2 and D3 is not mentioned in the description, nor are these documents identified therein.
- 10.3 The features of the claims are not provided with reference signs placed in parentheses (Rule 6.2(b) PCT).
- 10.4 Furthermore, at page 40, last paragraph, the description contains general statements that the extent of protection may be expanded in some vague and not precisely defined way. Such general statements shall be deleted as contrary to Article 6 PCT, cf. also PCT Preliminary Examination Guidelines, C-III, 4.3a.

Re Item VIII

Certain observations on the international application

- 11 Present independent claim 1 is not clear and as such it does not fulfill the requirements of Article 6 PCT for the following reasons.
- 11.1 The wording of present claim 1 is too broad and vague and it does not enable the person skilled in the art to carry out the invention without any further inventive effort, which is against Article 6 PCT. It is the opinion of this Authority that the person skilled in the art would be unable. on the basis of the information given in the application as filed, to extend the particular teaching of the description to the whole of the fields claimed by using routine methods of experimentation or analysis. Therefore, the wording of present claim 1 should be amended in order to properly limit the extent of the subject matter claimed in accordance with the subject matter as disclosed in the application as a whole.
- 12 Dependent claim 2 is also vague and unclear and therefore it does not meet the requirements of Article 6 PCT, the reasons therefor being the following.

- 12.1 The wording "comprising the step of employing discretized wavelet transform analysis to process the cardiac waveform" used in claim 2 is vague and unclear and leaves the reader in doubt as to the meaning of the technical features to which it refers, thereby rendering the definition of the subject matter of said claim unclear (Article 6 PCT).
- 13 Dependent claim 3 is also vague and unclear and therefore it does not meet the requirements of Article 6 PCT, for the same reason already mentioned in paragraph 12.1 of this International Preliminary Examination Report and for the following additional reason.
- 13.1 The wording "discretized continuous" used in claim 3 does not have a clear technical meaning and as such it leaves the reader in doubt as to the meaning of the technical features to which it refers, thereby rendering the definition of the subject matter of said claim unclear (Article 6 PCT). This is also so because a wavelet basis either is discrete or is continuous; moreover it appears from the description that the method of the application is specified in base of continuous Morlet wavelet basis functions.
- Dependent claim 7 is also vague and unclear and therefore it does not meet the 14 requirements of Article 6 PCT, the reasons therefor are the following.
- 14.1 The wording "derive the cardiac signal" used in claim 7 is unclear and as such it leaves the reader in doubt as to the meaning of the technical features to which it refers, thereby rendering the definition of the subject matter of said claim unclear for the following reasons.
 - The wording of present claims 1 to 6 does not specify whether the named "cardiac signal" is an analogical signal or it is a digitized signal, while from the wording of dependent claim 7 it appears that what the applicant meant was a digitized cardiac signal. This fact creates in the reader a state of uncertainty as to the extent of the subject matter claimed, which is against the provisions of Article 6 Pct.

EXAMINATION REPORT - SEPARATE SHEET

- Dependent claims 10 and 11 are also vague and unclear and as such they do not 15 fulfill the provisions of clarity of Article 6 PCT for the following reasons.
- 15.1 The wordings "coherent structures" and "for clinical use" used in claim 10 and 11 respectively do not have a clear technical meaning and as such they leave the reader in doubt as to the meaning of the technical features to which it refers, thereby rendering the definition of the subject matter of said claim unclear (Article 6 PCT).
- Claim 15 does not meet the requirements of Article 6 PCT in that the matter 16 claimed is not clearly defined. The claim attempts to define the subject matter in terms of the result to be achieved which merely amounts to a statement of the underlying problem. The technical features necessary for achieving this result should be added.
- Independent claim 16 does not meet the requirements of Article 6 PCT in that the 17 matter claimed is not clearly defined. The overall wording of claim 16 does not provide the required intelligibility in order to be understood by the average skilled person. It is necessary to redraft the claim using a more understandable manner, in order to meet the requirements of Article 6 PCT (clarity in the sense of intelligibility).
- 18 Present independent claim 33 is not clear and as such it does not fulfill the requirements of Article 6 PCT for the same reasons already mentioned in paragraph 11.1 of this International Preliminary Examination Report.
- 19 Independent claim 34 and dependent claims 35 to 37 are also vague and unclear and as such they do not fulfill the requirements of Article 6 PCT for the following reasons.
- 19.1 Claims 34 to 37 do not meet the requirements of Article 6 PCT in that the matter

INTERNATIONAL PRELIMINARY



International application No. PCT/GB00/01675

EXAMINATION REPORT - SEPARATE SHEET

claimed is not clearly defined. The functional statements included in the wording of present claims 34 to 37 do not enable the skilled person to determine which technical features are necessary to perform the stated functions.

Independent claim 38 does not meet the requirements of Article 6 PCT in that the 20 matter claimed is not clearly defined for the same reasons as above mentioned in paragraph 17 of this International Preliminary Examination Report.

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REQUEST

The undersigned requests that the present international application be processed according to the Patent Cooperation Treaty.

For re-	œiving Of GB O	fice use onl	y —	6 7	5
International Application 1	No.				
International Filing Date	02-	OS-		2000	
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Box No. I T	ITLE OF IN	VENTION				-	-
	"Method	of Analysis of	Medical Signal	s"			
Box No. II A	PPLICANT			_			
Name and address designation. The address indicated of residence is indicated to the second s	address must i l in this Box is t	include postal coc the applicant's St	iven name; for a le and name of cou ate (that is, country	legal entity, full of ntry. The country of residence if no	ficial of the State	This p	erson is also inventor.
The Court of N	•	ersity				Telephone No.	
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Box No. III F	URTHER A	PPLICANT(S) A	ND/OR (FURTI	IER) INVENTO	R(S)		
ADDISON, Pau 58 Buckstone (EDINBURGH EH10 6UR United Kingdor	ul Stanley Crook n		iven name; for a le and name of cou ute (that is, country,			applica invento is marka	is: ant only nt and inventor or only (If this check-box ed, do not fill in below.)
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This person is ap for the purposes		all designated States	all designated the United St	States except ates of America		United States America only	the States indicated in the Supplemental Box
Further app	licants and/or	(further) invento	ers are indicated or	n a continuation sl	heet.	. –	
Box No. IV A	GENT OR C	OMMON REPI	RESENTATIVE;	OR ADDRESS	FOR C	ORRESPONDI	ENCE
The person identi- of the applicant(s)	ied below is l before the co	hereby/has been a ompetent Internat	appointed to act or ional Authorities a	behalf as:	X ag	gent	common representative
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Sheet No. 2

Continuation of Box No. III FURTHER APPLICANT(S) AND/OR (FURTHER) INVENTOR(S)							
If none of the following sub-boxes is used,	this sheet should not be included in the request.						
Name and address: (Family name followed by given name; for a designation. The addressmust include postal code and name of con address indicated in this Box is the applicant's State (that is, country of residence is indicated below.) WATSON, James Nicholas 34 Fowler Terrace EDINBURGH EH11 1DA United Kingdom	legal entity, full official unity. The country of the poly of residence if no State This person is: applicant only applicant and inventor inventor only (If this check-box is marked, do not fill in below.)						
State (that is, country) of nationality: United Kingdom	State (that is, country) of residence: United Kingdom						
This person is applicant for the purposes of: all designated the United States all designated the United States	d States except the United States the States indicated in the Supplemental Box						
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This person is applicant all designated the United States all designated for the purposes of: Name and address: (Family name followed by given name; for a last designation. The address must include postal code and name of cour address indicated in this Box is the applicant's State (that is, country) of residence is indicated below.)	States except the United States the States indicated in of America only the Supplemental Box egal entity, full official ciry. The country of the of residence if no State This person is: applicant only applicant and inventor inventor only (If this check-box is marked, do not fill in below.)						
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Further applicants and/or (further) inventors are indicated or	another continuation sheet.						





Sheet No. 3.....

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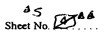
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1 "Method of Analysis of Medical Signals"

2

- 3 This invention relates to a method of analysis of
- 4 medical signals, and in particular to a method of
- 5 decomposition of cardiac signals using wavelet
- 6 transform analysis. Specifically the invention relates
- 7 to an improved method of resuscitation of patients in
- 8 cardiac arrest.

- 10 In the UK, coronary heart disease is the second
- 11 greatest contributor to deaths of people under 75. The
- 12 social and economic consequences of these death rates





The current survivability rates of are enormous. 1 patients after sudden cardiac failure are around 1:10. 2 3 Ventricular tachyarrhythmias, specifically ventricular 4 fibrillation (VF), are the primary arrhythmic events in 5 cases of sudden cardiac death. Administration of 6 prompt therapy to a patient presenting with such 7 symptoms can however lead to their successful 8 resuscitation. Until recently, the only indicators of 9 likelihood of survival of a patient to hospital 10 discharge were traditional variables such as emergency 11 service response time or bystander cardio-pulmonary 12 13 resuscitation (CPR). 14 In most cardiac complaints, analysis of a surface 15 electrocardiogram (EKG) of the presenting patient is a 16 rich source of information. However, until recently, a 17 surface EKG recorded during VF and any subsequent 18 19 medical intervention to defibrillate, was thought merely to present unstructured electrical activity, and 20 21 not to provide useful information. 22 The first attempts to derive prognostic information 23_ from EKGs of the heart in VF focussed on the importance 24 25 of the amplitude of the waveform defined using peak-totrough differences in the EKG voltage, measured as 26 either the greatest deflection occurring in a 27 predefined time slot, or as the average peak-to-trough 28 29 voltage measured over a given time interval. been shown that the VF amplitude is inversely related 30 to time elapsed since collapse, is a crude predictor of 31 defibrillation outcome, and is a better indicator of

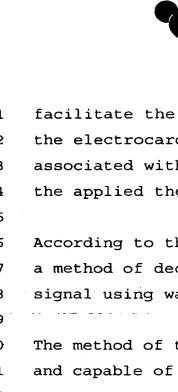


1	survival to hospital discharge than the traditional
2	variables described above.
3	
4	However, recording the VF amplitude accurately is
5	significantly problematical. The EKG voltage amplitude
6	measured during VF is dependent on the direction of the
7	main fibrillation vector and is influenced by a variety
8	of factors including patient chest shape; electrode
9	size; electrode location; and skin/electrode interface
10	resistance. This number of variables makes this
11	amplitude measure both unreliable and inaccurate. That
12	is, although the amplitude of the waveform of an EKG
13 .	recorded during VF is now recognised to be a crude
14	predictor of the likely outcome of resuscitation of a
15	patient in VF, it is not a reproducible marker of
16	sensitivity to defibrillation, and lacks clinical
17	usefulness.
18	
19	In a further development, it is also known to use Fast-
20	Fourier based transforms to generate a frequency
21	spectrum of an EKG in VF to analyse the signal. The
22	median frequency (MF) divides the area under the
23	spectrum into two equal parts. Since this plot is
24	derived from information in both the voltage and time
25	domains, external variables such as lead placement have
26	less effect on the results than the method of observing
27	the amplitude. However, CPR produces artefacts in the
28	recorded EKG signal and, since pausing CPR merely to
29	obtain an EKG signal free of artefacts is likely to
30	compromise resuscitation, these artefacts are
31	necessarily included in this frequency measure, and
32	detract from its usefulness.

•

4

1 2 Thus the results of such signal analysis show some 3 correlation with the likely outcome of resuscitation, but again lack sufficient sensitivity and specificity 4 5 for clinical use. That is, this form of analysis has 6 the disadvantage that, since the Fourier spectrum 7 contains only globally averaged information, specific 8 features in the signal are lost. 9 A method of accurate analysis of a surface EKG waveform 10 11 recorded during VF would therefore be useful in 12 understanding the pathophysiological processes in 13 sudden cardiac death, and thus to produce a model for 14 use: 15 16 in predicting the efficacy of therapy in individual 17 cases; and 18 in determining the selection of the preferred course of 19 20 primary, and alternative or adjunct therapies thus 21 providing a means for individually tailored therapy for 22 the specific patient needs 23 24 to improve the success rate of resuscitation of 25 patients presenting in VF. 26 27 Atrial fibrillation (AF) is a common cardiac arrhythmia 28 in older people. Atrial fibrillation can be stopped by giving an electric shock to the patient under general 29 30 anaesthetic (cardioversion). However, many patient 31 return to an AF rhythm soon after treatment. The technology detailed here may also provide a tool to 32



facilitate the clinical evaluation of AF exhibited in 1

the electrocardiogram (EKG) so reducing the risk

3 associated with general anaesthetic in patients where

the applied therapy is likely to prove ineffective.

According to the present invention there is provided 6

a method of decomposition of waveforms in a cardiac 7

8 signal using wavelet transform analysis.

10 The method of the invention is non-invasive, accurate,

11 and capable of delivering real-time information.

12

Preferably said method employs discretized wavelet 13

transform analysis to process the EKG. 14

15

16 Preferably said method employs discretized continuous

17 wavelet transform analysis to process the EKG.

18

Preferably said method comprises the steps of deriving 19

20 the wavelet energy surfaces of an EKG signal; and

21 plotting said wavelet energy surfaces against a

22 location parameter b, and a scale parameter. The scale

23 parameter may be dilation a or band pass frequency f_{bpc} .

24

25 The method initially comprises the steps of connecting

26 electrodes to the presenting patient; and sampling the

27 analogue input signal to derive the cardiac signal.

28

29 Typically said method comprises the step of visually

30 displaying the cardiac signal.



Said method may display the distribution of energies 1 within the cardiac signal. Said method may display 2 coherent structures within the cardiac signal. 3 4 Said display may be by means of a contour plot. 5 display may be by means of a surface plot. 6 said method provides means to visualise the signal in 7 real-time for clinical use. 8 9 Preferably said method is applicable in the analysis of 10 an EKG in ventricular fibrillation. 11 12 Said method may be applicable in the analysis of an EKG 13 in ventricular fibrillation after the commencement of 14 15 cardio-pulmonary resuscitation (CPR). 16 The method may include the step of disassociating the 17 component features of the temporal trace of a recorded 18 Additionally or alternatively said method may 19 include the step of temporal filtering of an EKG signal 20 of a heart which is subject to CPR to disassociate the 21 22 CPR signal from the heart signal. 23 Typically said method provides measurable 24 characteristics for the estimation of the health of a 25 26 heart in VF. Said method may provide measurable characteristics for the estimation of the health of a 27 Said me may provide Typically said method 28 heart in AF. provides measurable characteristics for the estimation

29

30 31 of the health of a heart.



The method may provide measurable characteristics for 1 2 the estimation of the time elapsed since the onset of a 3 cardiac incident. 4 Typically said method provides measurable 5 characteristics for the estimation of the health of a 6 heart after commencement in CPR. 7 8 Said method may provide a prediction for the outcome of a given therapeutic intervention and so aid the 10 11 clinical decision making process. 12 Said method may provide a basis for individual, patient 13 14 specific, protocols for therapeutic intervention. 15 16 The method may provide a guide to the optimal timing of defibrillation of a heart in VF. 17 18 Said method may include the step of constructing a 19 damage index for reference purposes. Construction of 20 said index might involve the development of a network 21 22 classifier from a library of recorded data. network classifier may comprise a neural network. Said 23 network classifier may comprise a wavelet network 24 25 classifier. 26 Application of the method of the invention represents a 27 significant advance in coronary care by providing a

significant advance in coronary care by providing a reliable predictor of the outcome of shocking a patient in VF. In addition, the development of an algorithm using the method of the invention gives the ability to predict shock outcome and to facilitate individual

8

patient therapy. The ability to provide patient 1 2 specific therapeutic intervention is a priority in the 3 advancement of currently applied medical protocols. That is, as discussed above, in certain instances, 5 after prolonged cardiac arrest preceding defibrillation 6 7 pharmacological measures or CPR can increase the chance 8 of successful resuscitation. Thus, employing the method to predict the outcome of shocking avoids futile 9 defibrillation attempts which can even harm the heart, 10 11 and can indicate the need for intervention, and 12 influence the selection of the preferred type of intervention, to optimise the metabolic state of the 13 heart prior to counter-shock. 14 15 16 The predictor algorithm developed using the method is 17 being tested using a new generation of defibrillation 18 devices that have the flexibility to allow easy prototyping of the new defibrillation algorithms. 19 20 21 According to a further aspect of the present invention 22 there is provided a method of decomposition of 23 waveforms in a cardiac signal using matching pursuit 24 algorithms. 25 26 According to a further aspect of the present invention 27 there is provided an apparatus for decomposition of 28 waveforms in a cardiac signal, said apparatus 29 comprising wavelet transform analysis means. 30 31 Said apparatus may include means to display the 32 distribution of energies within a waveform.

phonographic signals.

1	Said apparatus may include a monitor adapted to display
2	decomposed waveforms. Said apparatus may be adapted
3	for inclusion in an EKG apparatus.
4	
5	According to a further aspect of the present invention
6	there is provided defibrillation means adapted to
7	operate in response to a signal generated by comparison
8	of an EKG trace with decomposed waveform.
9	
10	That is, the invention preferably provides a method of
11	wavelet analysis of cardiac signals which provides
12	structural information about the heart - whether the
13	heart is healthy or not - and has significant
14	advantages over fast Fourier transforms.
15	
16	The invention may provide a display device in the form
17	of a scrologram that provides real-time visualisation
18	of a wavelet scalogram, showing the distribution of
19	energies and coherent structures within the signal for
20	use as guidance by a clinician.
21	
22	The invention may further provide a data analysis tool,
23	which assists in shock timing (atrial pulsing). That
24	is, the derived data may indicate the optimum time to
25	administer shock to the heart. The invention may
26	provide a damage index, preferably in the form of an
27	artificial neural network.
28	
29	Preferably the invention provides dissociation of the
30	component features of a temporal trace of a cardiac
31	signal, which may for example be CPR, AF, or cardio-



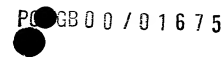
Ţ	Embodiments of the invention will now be described by
2	way of example only and with reference to the
3	accompanying drawings in which:
4	
5	
6	Figure la is a Mexican hat wavelet;
7	
8	Figure 1b is the real part of a complex Morlet
9	wavelet;
10	
11	Figure 2a is a schematic plot showing the dilation
12	of a continuous wavelet;
13	
14	Figure 2b is a schematic plot showing the
15	translation of a continuous wavelet;
16	
17	Figures 3a to Figure 3e are the plots of the
18	'investigation' of a sinusoidal signal by Mexican
19	hat wavelets of various sizes, showing the effect
20	of translation of the wavelet along the signal
21	(change in b), and dilation of the wavelet (change
22	in a);
23	
24	Figure 4a is the plot of five cycles of a sine
25	wave of period P;
26	
27	Figure 4b is the contour plot of $T(a,b)$ against a
28	and b for the sine wave of Figure 4a;
29	
30	Figure 4c is the isometric surface plot of $T(a,b)$
31	against a and b for the sine wave of Figure 4a;
32	



1	Figure 5a is the plot of a combination of two sine
2	waves of period P1, and P2, where P1 = 5P2;
3	
4	Figure 5b is the contour plot of $T(a,b)$ against a
5	and b for the sine wave of Figure 5a;
6	
7	Figure 5c is the isometric surface plot of $T(a,b)$
8 .	against a and b for the sine wave of Figure 5a;
9	
10	Figure 6a is an EKG trace of a pig heart in sinus
11	rhythm;
12	
13	Figure 6b is a 2D energy scalogram associated with
14	the EKG trace of Figure 6a;
15	
16	Figure 6c is a 3D energy scalogram associated with
17	the EKG trace of Figure 6a;
18	
19	Figures 6d, 6e, 6f and 6g are the energy surface
20	plots from four segments of an EKG signal
21	subsequent to the onset of VF, showing the three
22	dominant ridges A, B, and C appearing in the
23	transform surface, and showing in Figure 6g the
24	onset of CPR after five minutes, associated with a
25	gradual increase in passband frequency of the
26	ridges A,B, and C;
27	
28	Figure 7a is an energy scalogram for a pig heart
29	for the first seven minutes of ventricular
30	fibrillation, indicating the initiation of CPR
31	after five minutes;
32	



1	Figure 7b is a schematic diagram of the salient
2	features of the scalogram of Figure 7a;
3	
4	Figure 7c is the smoothed plot of energy at the
5	8Hz level in the scalogram of Figure 7a against
6	time;
7	
8	Figure 8a is a typical segment of an EKG trace of
9	a pig heart in VF;
10	
11	Figures 8b, 8c, and 8d are the energy scalograms
12	associated with the trace of Figure 8a;
13	
14	Figure 9 is a screen shot of a real time viewer
15	which shows the collected EKG data with its
16	associated wavelet energy display in the form of
17	its energy scalogram, where windows scroll to the
18	right;
19	
20	Figure 10a is a 7 second trace of human ECG
21	showing a shock event;
22	·
23	Figure 10b is a scalogram corresponding to the
24	trace of Figure 10a;
25	
26	Figure 11a shows the proportion of energy in
27	scalograms for 120 results (60 ROSC, and 60
28	asystole) at 1.9 Hz after shocking;
29	
30	Figure 11b shows the proportion of energy in
31	scalograms for 120 results (60 ROSC, and 60
32	asystole) at 9.3 Hz after shocking;



1 2 Figure 12a is a schematic representation of 3 overlapping signal segments used in a neural 4 network test study; 5 6 Figure 12b shows the weights attributed by the 7 Kohonen network to the 30 frequency levels used in 8 the scalogram; 9 Figure 13a is an aorta pressure trace; 10 11 12 Figure 13b shows the EKG for the same time period as the trace of Figure 13a; and 13 14 15 Figure 13c is the scalogram associated with the 16 trace of Figure 13a derived from the Morlet 17 wavelet; 18 Figure 13d is a detail of the phase part of 19 20 scalogram Figure 13c; 21 22 Figure 13e is the scalogram associated with the 23 trace of Figure 13a derived from the Mexican hat 24 wavelet; and 25 26 Figure 13f demonstrates the correlation of aorta 27 pressure pulse position with lines of zero phase; 28 29 Figures 14a is the plot of an EKG trace. Figure 30 14b is its associated phase at around 1.5Hz. 31 Figure 14c is its energy scalogram. The 32 correlation of zero phase at this lower frequency

and high frequency (low dilation) peaks is thus 1 2 illustrated. 3 Figure 15a shows a 2 second segment of EKG taken 4 from a patient with atrial fibrillation (AF). 5 Figure 15b shows the wavelet scalogram plot 6 associated with this EKG. Figure 15c shows the 7 corresponding modulus maxima of the scalogram of 8 Figure 15b. 9 10 Figure 15d contains a 7 second segment of EKG 11 12 exhibiting AF. Figure 15e is a trace of EKG temporal components with small amplitude. Figure 13 14 15f shows the larger magnitude components i.e. the ORS and T waves. 15 16 Figure 15g is a plot of a two second 'blow up' of 17 part of the signal of Figure 15d; Figure 15h is a 18 19 plot of a two second 'blow up' of part of the signal of Figure 15e; and Figure 15i is a plot of 20 a two second 'blow up' of part of the signal of 21 22 Figure 15f. 23

24 Referring to the Figures, the present method employs 25 the use of a wavelet transform to analyse a cardiac

26 signal.

27

28 The method involves the decomposition of the signal.

This decomposition is accomplished by utilising wavelet 29

30 transforms to decompose the signal in wavelet space.



A key distinction between the Fourier analysis of an

2 EKG signal and its analysis by means of a wavelet function is that, whilst the Fourier transform employs 3 a sinusoid function, a wavelet function is localised in 4 time. 5 6 7 The methodology for such decomposition may include discretized continuous wavelet transforms, orthonormal 8 wavelet transforms of decimated construction, non-9 decimated wavelet transforms, wavelet packet transforms 10 and matching pursuit algorithms. 11 12 13 Signal processing employing wavelet transform analysis 14 allows simultaneous elucidation of both spectral and temporal information carried within a signal. 15 processing can employ either continuous or discrete 16 The choice of wavelet transform used for a 17 transforms. particular signal processing application depends on 18 factors such as speed of computation necessary, the 19 shape of signal specific features, the frequency 20 resolution required, and the statistical analysis to be 21 22 performed. Andrews and the second of the 23 24 The preferred method employs the discretized continuous 25 transform as it provides high resolution in wavelet space at lower frequencies. 26

27

1

-28 This method thus employs the use of a discretized continuous wavelet transform to analyse a cardiac 29

30 signal.

- 1 In particular, this method employs a wavelet transform
- 2 as an interrogation tool for EKG signals of ventricular

16

3 fibrillation.

4

- 5 A variety of wavelet functions are available, and the
- 6 most appropriate is selected to analyse the signal to
- 7 be investigated.

8

- 9 The wavelet transform of a continuous time signal,
- 10 x(t), is defined as:

11

12
$$T(a,b) = \frac{1}{w(a)} \int_{-\infty}^{\infty} x(t) \overline{g} \left(\frac{t-b}{a} \right) dt$$

equation 1

13

- 14 where g(t-b)/a) is the analysing wavelet function and
- 15 ' denotes complex conjugate. w(a) is a scaling
- 16 function usually of the form $w(a) = a^n$ where n is usually
- 17 1 or 0.5, and x(t), in this application, is the single
- 18 channel surface EKG time signal. The transform
- 19 coefficients T(a,b) are found for both specific
- 20 locations on the signal, b, and for specific wavelet
- 21___dilations, a.__T(a,b)_is_plotted_against_a_and_b_in_____
- 22 either a surface or contour plot.

- 24 While other wavelet types may be employed the wavelets
- 25 mainly used in this method are: the Mexican hat wavelet
- 26 and the Morlet wavelet, examples of which are shown in
- 27 Figure 1.





- 1 The wavelet can translate along the signal (change in
- 2 b) and dilate (change in a). This is shown
- 3 schematically in Figure 2 using a Mexican hat wavelet.
- 4 Figure 3 illustrates the way in which a sinusoidal
- 5 signal can be 'investigated' at various locations by
- 6 Mexican hat wavelets of various sizes. The numerical
- 7 value of the convolution (equation 1) depends upon both
- 8 the location and dilation of the wavelet with respect
- 9 to the signal.
- 10 Figure 3a shows a wavelet of similar 'size' to the
- 11 sinusoidal waves superimposed on the signal at a b
- 12 location which produces a reasonable matching of the
- 13 wavelet and signal locally. From the Figure it is
- 14 apparent that there is a high correlation between the
- 15 signal and wavelet at this a scale and b location.
- 16 Here, the cross correlation of the signal with the
- 17 wavelet produces a large positive number T(a,b).
- 18 Figures 3b and 3c show details of the wavelet transform
- 19 of a signal using a wavelet of approximately the same
- shape and size as the signal in the vicinity of b.
- 21 Figure 3b-shows a wavelet of similar scale to the
- 22 sinusoidal waveform located at maximum negative
- 23 correlation. This produces a large negative T(a,b)
- 24 value. Figure 3c shows a wavelet of similar scale to
- 25 the sinusoidal waveform located at a position on the
- 26 time axis where near zero values of T(a,b) are
- 27 realised. Figure 3d shows the effect on the transform
- 28 of using the smaller a scale. It can be seen from the
- 29 plot that the positive and negative parts of the
- 30 wavelet are all in the vicinity of approximately the



- 1 same part of the signal, producing a value of T(a,b)
- 2 near zero. Figure 3e shows that the same thing happens
- 3 when using a much larger wavelet, since the wavelet
- 4 transform now covers various positive and negative
- 5 repeating parts of the signal, again producing a near
- 6 zero value of T(a,b).

- 8 Wavelet transforms are not usually computed at
- 9 arbitrary dilations for isolated locations in the
- 10 signal, but rather over a range of a and b. A plot of
- 11 T(a,b) versus a and b for sinusoidal data using the
- 12 Mexican hat wavelet is shown in Figure 4. Two methods
- are then employed to plot T(a,b), namely a contour plot
- or scalogram as shown in Figure 4b, and a surface plot
- 15 as shown in Figure 4c. At small and large values of a,
- 16 the near zero values of T(a,b) are evident from the
- 17 plots, but at values of a of the order of one quarter
- 18 of the wavelength of the sinusoid large undulations in
- 19 T(a,b) correlate with the sinusoidal forms of the
- 20 signal.

- 22 Figure 5a shows two superpositioned sinusoidal
- 23 waveforms, the first with period P1, the second with
- 24 period P2. P1 = 5P2. Figures 5b and 5c, the transform
- 25 plots of the superimposed waveforms clearly show the
- 26 two periodic waveforms in the signal at scales of one
- 27 quarter of each period. Thus, Figure 5 clearly
- 28 demonstrates the ability of the continuous wavelet
- 29 transform to decompose the signal into its separate





- 1 frequency components. That is, this transform
- 2 'unfolds' the signal to show its constituent waveforms.
- 3 The contribution to the signal energy at a specific a
- 4 scale and b location is proportional to the two-
- 5 dimensional wavelet energy density function which is,
- 6 in turn, proportional to the modulus of T(a,b).

- 8 The method of the present invention thus involves the
- 9 display of the transform as a contour plot. That is,
- 10 the method is used to present information derived from
- 11 an EKG trace of the heart in VF as a scalogram. The
- 12 preferred form of presenting the information is as an
- 13 energy scalogram, which presents the results as a plot
- 14 showing the log of the wavelet energy coefficients,
- 15 against the log of the bandpass centre frequency, f_{bpc} ,
- 16 of the wavelets for each time increment. The bandpass
- 17 centre frequency is proportional to the reciprocal of
- 18 the dilation value, a. This plot highlights small
- 19 changes in amplitude over the scales of interest. The
- 20 transform copes with repeating features in time with
- 21 shifting-phase,-making-it-appropriate for-real time —
- 22 applications such as this.

- 24 That is, by performing continuous wavelet transform
- 25 analysis on the ECG in VF, and then by producing an
- 26 energy scalogram of the results, it is possible to
- 27 unfold the signal in such a way that a previously
- 28 hidden structure is apparent, in contrast to the
- 29 apparently disorganised VF signal.





- 1 The method then includes quantifying the wavelet
- 2 decomposition. This wavelet decomposition provides
- 3 both qualitative visual and measurable features of the
- 4 EKG in wavelet space.

- 6 In practice, surface EKG tracings, recorded as soon as
- 7 possible after the onset of VF, are analysed.

8

- 9 As a demonstration of the efficacy of the method, in an
- 10 example of an experimental procedure utilising this
- 11 method of analysis employing wavelet techniques, VF was
- 12 induced in anaesthetised pigs via a pacemaker probe,
- 13 using a 90V impulse at 60 Hz. All of the pigs remained
- in VF, untreated for a period of either 3 or 5 minutes.
- 15 After this time, CPR commenced. The surface EKG
- 16 (standard lead II) was recorded using needle
- 17 electrodes. The EKG was sampled at 300 Hz using a 12-
- 18 bit A to D converter. The method of the present
- 19 invention was then performed using 32 EKG tracings
- 20 recorded immediately after the onset of VF.

21

- 22 Figure 6a represents 4 beats of a pig heart in sinus
- 23 rhythm. Figures 6b and 6c shows the wavelet transform
- 24 of the signal displayed in two and three dimensions
- 25 respectively.

- 27 The QRS complex of the waveform is evident from the
- 28 conical structures in Figure 6b converging to the high
- 29 frequency components of the RS spike. The P and T
- 30 waves are also labelled in the plot. The 3D landscape
- 31 plot of Figure 6c shows the morphology of the signal in



- 1 wavelet space. In Figures 6b and 6c the continuous
- 2 horizontal band (X) is associated with a frequency of
- 3 1.7 Hz, the beat frequency of the sinus rhythm. The
- 4 second band (Y) occurs at a frequency of approximately
- 5 5.1 Hz, corresponding to the separation of the P-QRS-T
- 6 components in time. At higher frequencies the P, QRS
- 7 and T components are individually resolved according to
- 8 their frequency makeup and temporal location.

- 10 Figures 6d to 6g show the energy surfaces for four
- 11 segments of EKG signal subsequent to the onset of VF,
- 12 namely: (6d) 0-60 s; (6e) 60-100 s; (6f) 210-240 s;
- 13 and (6g) 260-360 s.

14

- 15 The morphology of the VF signal in wavelet space can be
- 16 seen from the Figures to contain underlying features
- 17 within a more complex surface topography. The most
- 18 significant features are the dominant ridges that
- 19 appear in the transform surface through time.

20

- 21 Figure 6f shows these ridges quite clearly. A high-
- 22 energy ridge can be observed at around 10 Hz and two
- 23 lower energy bands can be observed at lower
- 24 frequencies. These three ridges are labelled A, B and
- 25 C, respectively, in the plot. Other ridges are also
- 26 present within the scalogram.

- 28 The energy surface in Figure 6g contains the onset of
- 29 CPR after 5 min of untreated VF. The institution of
- 30 CPR is associated with a gradual increase in the
- 31 passband frequencies of ridges A, B and C. This change
- 32 in the composition of the VF signal reflects electrical



changes in the fibrillating myocardium associated with 1 This is because CPR produces 2 the onset of CPR. 3 antegrade myocardial blood flow and thus improves the metabolic state of the tissues, temporarily reversing 4 the otherwise progressive decline in high band pass 5 frequency components of the EKG wavelet decomposition. 6 7 Figure 8a is a typical segment of an EKG trace of a pig 8 heart in VF; Figures 8b, 8c, and 8d are the energy 9 scalograms associated with the trace of Figure 8a. As 10 clearly illustrated by these diagrams the principle 11 dilation (band pass centre frequency) component of the 12 scalogram is approximately 10Hz. However, using said 13 method it is also apparent that this component is not 14 constant. It 'pulses' with a degree of regularity. This 15 structure is previously unreported. 16 17 18 Figure 9 shows similar 'pulsing' in another porcine EKG signal. However, the structure is so pronounced that 19 high energy, high frequency, intermittent components 20 can be observed. These components have an occurrence 21 22 frequency of the order of the original sinus rhythm: approximately 1.7Hz.

24

23

Figure 10a is a human EKG signal segment containing a 25 shock event. Figure 10b is the corresponding wavelet 26 scalogram. It is apparent from the scalogram of Figure 27 10b that both high frequency spiking and an 28 29 intermittent high-energy region are present in the 30 vicinity of 10 Hz and also above 10Hz.



- 1 The high frequency spiking is unique to the method of
- 2 the present invention and is not visible using
- 3 conventional Fourier techniques. The rich structure
- 4 made visible within the EKG by the wavelet transform
- 5 method is evident in the scalogram.
- 6 It is clearly seen from the Figures that applying the
- 7 wavelet transform to an EKG signal of VF demonstrates
- 8 that this signal is a rich source of valuable
- 9 information. That is, it produces a display showing
- 10 real time visualisation of the distribution of energies
- 11 and coherent structures within the signal for use by a
- 12 clinician in the selection of treatment strategies.
- 13 Using this method of analysis it is feasible to obtain
- 14 real-time visual display of the EKG frequency
- 15 characteristics in the wavelet domain during
- 16 resuscitation. The scalogram produced provides
- 17 information about the myocardium that is not available
- 18 from a standard single channel surface EKG.

- 20 The wavelet scalogram decomposition can be displayed as
- 21 a real time scrolling window, as shown in Figure 9.
- 22 This window is useful as an aid for clinical decision
- 23 making. It can be used as a stand-alone tool, or as
- 24 basis for on-line statistical analysis of the current
- 25 state of a heart.

- 27 To produce the window, a MATLAB TM R11 application is
- 28 used. Each EKG sample taken results in the updating of
- 29 a FIFO (First In First Out) buffer, and the EKG plot of
- 30 Figure 9a. The scalogram of Figure 9b is then shifted



to the right and clipped before the 'missing' new right 1 2 hand data is calculated, using conventional matrix algebra, and filled. 3 4 This results in the two scrolling windows of Figure 9. 5 6 The exponential ramp in the bottom right corner shows 7 the compact support of the wavelet utilised at the given scale. 8 9 10 Higher resolution scalograms are achieved through implementation on higher specification machines, 11 12 purpose built hardware, or application specific 13 software with coding using a lower level programming 14 language, such as C++. 15 16 CPR produces artefacts in the EKG signal. Additionally, 17 this method delivers information the value of which is 18 not degraded once the CPR artefacts are filtered from the EKG signal. 19 20 21 From examination of the scalograms shown in Figures 6q, 22 7a and 7b it can be seen that the VF signature and the 23 signature of the CPR artefacts occupy distinct areas of 24 the scalogram, which permits their separation. 25 26 Known techniques such as the Modulus maxima method are 27 now available to reduce the non-zero data points in the

This method reduces the topography

of the scalogram surface to a series of ridges, thereby

28

29

wavelet scalogram.



considerably reducing the amount of data required to 1 2 represent the signal in the wavelet space. 3 4 The modulus maxima obtained from a bandlimited signal with a wavelet of finite compact support in the 5 frequency domain defines a complete and stable signal 6 7 representation. 8 In this method, temporal filtering of the original EKG 9 signal to disassociate the CPR signature from the heart 10 signal can either be done directly, using the wavelet 11 energy scalograms, or indirectly through modulus maxima 12 This allows the heart to be monitored 13 techniques. without necessitating cessation of CPR to allow rhythm 14 15 recognition. 16 Further to the above, the method may also be applied to 17 patients suffering form atrial fibrillation (AF) as a 18 means of disassociating the prevalent QRS and T waves 19 from the remainder of the signal. 20 21 22 Wavelet decomposition of the ECG signal is performed 23 using an appropriate wavelet function. The modulus maxima technique is used to encapsulate the scalogram 24 information in a series of ridges. Filtering of the 25 signal is then undertaken using the modulus maxima 26 27 information and through reconstruction the clinically 28 useful information is isolated from the signal . 29 Specifically, Figure 15a shows the wavelet transform 30 decomposition of a 2 second segment of ECG taken from a 31 patient with atrial fibrillation. Below the ECG trace 32



maxima of the scalogram is plotted below the scalogram.

1 is a wavelet scalogram plot. The corresponding modulus

2

For example, Figure 15d contains a 7 second segment of 4 ECG exhibiting AF. The signal has been partitioned 5 6 using a modulus maxima ridge following algorithm. The 7 modulus maxima ridges have been separated into large 8 and small scale features by thresholding the signal at 9 a predetermined wavelet scale. A blow up of part of the 10 signal is given in the lower three plots in the figure: Figures 15g, 15h and 15i. The middle of these plots 11 12 contains the partitioned signal with the QRS complex and T wave filtered out revealing regular, coherent 13 14 features that appear at a frequency of approximately 400 beats per minute, typical of AF. The lower plot 15 contains the partition with the filtered out QRS and T 16 17 waves. Although, a relatively simple modulus maxima 18 technique was used in this pilot study whereby the 19 modulus maxima lines were simple partitioned into two 20 subsets, the ability of the technique to separate the 21 signal into QRS and T waves and underlying atrial 22 activity is evident from the results. It is known that 23 the decay in amplitude of a modulus maxima 24 corresponding to a signal feature can be a function of 25 the scale of the wavelet. It is possible to use this 26 property to separate the ridge coefficients into a 27 noisy and coherent part. In this way, 28 differentiation of the modulus maxima information can 29 be implemented within a more sophisticated algorithm. 30 This will facilitate the further separation

background noise, QRS and T waves, and atrial activity.



1 This method thus facilitates useful interpretation of

- 2 previously unintelligible EKG signals.
- 3 In patients presenting with uncoordinated rapid
- 4 electric activity of the ventricle of heart, known as
- 5 ventricular fibrillation (VF), there is no effective
- 6 pulse and myocardial blood flow ceases. Even the
- 7 institution of optimal cardio-pulmonary resuscitation
- 8 (CPR) of the patient does not achieve more than 30% of
- 9 the normal cardiac output. Ischaemia during cardiac
- 10 arrest leads to a rapid depletion of myocardial high-
- 11 energy phosphates, deterioration of transmembrane
- 12 potentials, and disruption of intracellular calcium
- 13 balance. Paradoxically, the myocardium in VF has
- 14 supranormal metabolic demands. For this reason
- 15 resuscitation attempts become less likely to succeed
- 16 with the passage of time, and electrical defibrillating
- 17 shocks increasingly result in asystole or EMD.

- 19 After prolonged cardiac arrest, the use of
- 20 pharmacological measures or CPR before attempting
- 21 defibrillation may increase the chances of successful
- 22 resuscitation. This invention provides a robust and
- 23 reliable method of analysis of the state of the
- 24 myocardium in VF that prevents attempts to defibrillate
- 25 at times that are unlikely to be successful, or even
- 26 harmful to the heart. This method also provides an
- 27 indication of the best way in which to optimise the
- 28 metabolic state of the heart prior to counter-shock.





- 1 The method includes steps to establish a standard
- 2 against which to evaluate collected data in a
- 3 particular incidence.

- 5 The method further employs use of measurable signal
- 6 characteristics derived from the position and amplitude
- 7 of features in the scalogram to estimate both the
- 8 condition of the myocardium, and downtime of the
- 9 subject while in VF.

10

- 11 The method thus provides for optimal treatment of the
- 12 heart in VF, so fulfilling specific patient needs, by
- 13 therapeutic intervention, if appropriate.

14

- 15 An energy scalogram such as that shown in Figure 7
- 16 displays three distinct bands, labelled A, B, C. It is
- 17 possible to derive quantifiable measures using
- 18 correlations between the location and energy content of
- 19 the bands.

20

- 21 Band A of Figure 7b represents the dominant energy band
- 22 seen in the scalogram of Figure 7a, and corresponds to
- 23 the tachycardic beating of VF. However the scalogram
- 24 is much more informative in that it also shows, as
- 25 bands B and C, the behaviour of other frequency
- 26 components of the signal which were previously
- 27 unreported.

.28

- 29 Figure 7a shows a 2D energy scalogram. It includes the
- 30 first 5 minute period of VF, followed by a 2.5 minute
- 31 period of CPR. The onset of CPR is clearly identified
- 32 by the distinct horizontal dark band in the lower right



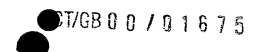
- 1 quadrant of the Figure. Over the first 5 minute
- 2 period, three bands, labelled A, B, C, can be clearly
- 3 seen in the scalograms. These bands correspond to the
- 4 ridges of Figures 6d to g. The increase in the
- 5 frequency components of these three bands after the
- 6 onset of CPR is evident in the plot. Bands B and C
- 7 follow trajectories similar to each other in the
- 8 scalogram, reducing in frequency over time. Band A,
- 9 however, moves independently of the other two.
- 10 Initially Band A increases, then it decreases to a
- 11 local minimum value at approximately 70s. Between 70
- 12 and 160s it increases relative to Bands B and C.
- 13 Finally, it decreases until the start of CPR after
- 14 300s. The same pattern was present in all 32 pig EKG
- 15 traces of the experiment.

- 17 Obvious increases in the passband frequency of all
- 18 three bands are observed in the scalogram after the
- 19 onset of CPR. For some of the signals studied this
- 20 increase in band C is masked by the dominant CPR band,
- 21 and thus cannot be seen in the scalogram.

22

23 Figure 7b provides a schematic diagram of the salient

- 24 features contained within the scalogram plots, where t0
- 25 is immediately after the onset of VF; t2 is the start
- of CPR; and t3 is the end of the analysis. Figure 7c
- 27 shows the relative proportion of energy contained in
- 28 the scalogram in the 5 to 12 Hz region through time.
- 29 There is an obvious decay in the relative energy
- 30 associated with this region which is associated with
- 31 the breakdown of co-ordinated activity in the heart.



- 1 The steps of the method of the present invention
- 2 described above establish that during the course of VF
- 3 there is a reduction in the proportion of energy within
- 4 the dominant frequency band indicated in Figure 7c.
- 5 This dominant frequency band, Band A in Figure 7a, is
- 6 demonstrated to be approximately 10 Hz for pig VF.

- 8 The energy within this band changes rapidly. This is
- 9 illustrated by the 'pulses' in Figures 8,9,10.

10

- 11 The Figures 6,7,8,9,10 show that applying the wavelet
- 12 transform to an EKG signal of VF demonstrates that this
- 13 signal is a rich source of valuable information.

14

- 15 The underlying hypothesis of the method of the present
- 16 invention is that the scalogram associated with an EKG
- 17 correlates to the state of the myocardium as it decays
- 18 subsequent to the onset of VF.

19

- 20 The method uses the information contained in the energy
- 21 scalogram associated with an EKG to predict the likely
- 22 success of clinical intervention, namely shocking.

23

- 24 It is therefore possible to develop a wavelet transform
- 25 based tool for the prediction of shock outcome during
- 26 ventricular fibrillation by:

27

- 28 1. collecting and collating data from sets of
- 29 archived EKGs recorded from humans in VF where
- 30 attempts to resuscitate by shocking were made; and

31

32 2. developing a classifier for reference purposes.

1 Figure 11 is a classification of the shock outcome in 2 either asystole or a rhythmic response using a 3 relatively simple statistical analysis. The experiment 4 yielding the results to compile these Figures involved 5 use of the lead II outputs of standard three lead EKGs of 120 patients in VF. Each trace is of three second 7 duration sampled at 100 Hz. Of these patients, 60 8 returned to sinus rhythm while the other 60 9 deteriorated to asystole, post shock. 10 11 Each trace was decomposed into an associated wavelet 12 transform from which its energy scalogram was 13 The volume under this surface was then 14 generated. normalised to render the results independent of signal 15 amplitude, but instead the result of the relative 16 wavelet constituents of the signals. The log of the 17 mean values at each dilation (band centre frequency) 18 for each was then recorded. Figures 11a and 11b show 19 the distribution of energies in a lower frequency band 20 (1.9 Hz) and at the 9.3 Hz band. Clearly, through 21 visual inspection, it is apparent that the proportion 22 of energies around the 10-Hz-band is higher-for 23 ---successful defibrillation attempts. 24 25 The method then extends to apply neural techniques to 26 analysis of wavelet pre-processed EKG signals. 27 28 A pilot study conducted to determine the feasibility of 29 using artificial neural techniques to provide a tool to 30 predict the outcome of defibrillation during VF used 31 eight human EKG trace segments containing shock events. 32

In these cases, the result of shocking was unequivocal

1

29

30 31

- four patients returned to VF, and four experienced 2 return of spontaneous circulation (ROSC). 3 4 The traces were transformed using the Morlet wavelet, 5 and energy scalograms containing thirty frequency 6 This was then split into eight 7 levels were produced. overlapping sections as shown in Figure 12a, each of 8 200 points (2/3 seconds duration). These 200 location 9 points were subsampled down to 50 to give eight 10 scalograms for each trace of 50 x 30 elements. 11 volume under the energy scalograms were normalised and 12 the patterns fed into a 'winner take all' Kohonen 13 network with two output units and built in conscience 14 15 (to avoid local minima). That is, the network was asked to group the 64 input patterns into two classes. 16 All but ten outputs were collectively classified 17 correctly giving a mean pattern error of 0.156 (against 18 0.5 average pattern error expected from random inputs). 19 20 Since this is a vector quantisation method (VQM) it was 21 possible to identify how the network differentiates the 22 patterns through inspection of its connective weights. 23 The weights from each location position across all 24 25 scales in the network are approximately the same, which means that there are no markers with which to 26 synchronise the different pre-processed traces. This 27 confirms that this neural network is too simple for 28

this purpose. That is the network is not equipped to

'consider' the relative phase of each input pattern.



+10%

Figure 12b shows the weights for the 'success' (ROSC) 1 and 'failure' (VF) to the output units from the first 2 two time slices across all scales. The weights 3 indicate the classes are differentiated by the 4 proportion of energy in the lower scales, which can be 5 seen when compared with Figure 11. 6 7 Although the above described method indicates the 8 slight drop in the dominant frequency expected, the 9 drop is very marginal which leads to the conclusion of 10 the lack of competence of previously proposed methods 11 as a defibrillation success predictor. 12 13 In summary, a library of human ECG data containing data 14 sets of human VF with attempts to resuscitate by 15 shocking is used as a database. This database is 16 extended to include data sets containing various 17 methods of shocking including, for example, biphasic 18 shocking. The biphasic shock waveform has resulted in 19 an increased proportion of successful defibrillation 20 attempts and is set to become the standard treatment 21 for cases of VF. 22 In one example, the recognised outcomes are defined by 24 trace components of the post-shock window lasting until 25 next shock (if present). If the ratio of the given 26 rhythm exceeds 10% of the total window length the 27 rhythms are prioritised according to the sequence: 28 29 Ratio Rhythm 30 Class 31

Pulse (SVR)

32



1	2	No pulse (EMD))	+10%	
2	3	Isoelectric (Asystole)	+10%	
3	4	VF		+10%	
4					
5					
6	Class 5 is the	class of VF pr	receding shocks w	here VF	
7	re-establishes itself within 5 seconds following the				
8	shock (i.e. no change). The VF in all the other				
9	classes were non-VF in this period.				
10					
11	Wavelet analysis of this information in accordance with				
12	the method of the invention is then performed to:				
13					
14	construct a wavelet visualisation of the signal -				
15	usually by plotting wavelet energy surfaces against the				
16	location parameter b and the inverse of the dilation				
17	parameter a;				
18					
19	provide measura	ble characteri	stics of the sign	nal for	
20	estimation of downtime of the patient;				
21					
22	provide measura	ble characteri	stics of the sign	nal for	
23	determining the	health of the	e heart post CPR;	and	
24					
25	to construct en	ergy scalogram	devised for the	method -	
26	which uses the energy density function and the				
27	reciprocal of the wavelet a scale for use as a				
28	predictor tool.				
29					
30	As described ab	ove it is poss	ible to use artif	icial	
31	neural network	based techniqu	es to develop suc	ch an	
32	indication of t	he state of my	ocardium. In the	e	





1 alternative, it is possible to classify the wavelet 2 scalogram through multilayered feedforward network 3 types. 4 The method may include the development of a modulus 5 maxima algorithm tool for the preprocessing of ECG 6 prior to its input into a neural network classifier. 7 8 9 Using this technique improves network performance whether this data is further encoded, or presented as a 10 whole, larger, sparse matrix as a pattern in the input 11 12 space. 13 This method therefore utilises the generalisation 14 properties of a feed forward multi-layer network to 15 predict the likelihood of defibrillation success from 16 the wavelet transform of the EKG traces. 17 layer network with its relatively simple dynamics, when 18 combined with wavelet pre-processing, has proved itself 19 20 a useful tool as a universal approximator. 21 The classes of multi-layer network types of use in this 22 23method are: 24

- Multi-layered feed forward (MLFF) neural networks 25 with back propagation training and monotonic 26 activation functions; and 27
- Radial Basis Neural Networks (RBNN) as have 28 previously been successfully applied to the denoising 29 of medical Doppler ultrasound signals with wavelet 30 31 preprocessing.



1 As described above, the method involves the 2 3 decomposition of EKG signals into a complete basis set defined by the wavelet shape and other parameters by 4 salient basis functions of a different basis set, 5 converged upon through regression techniques (sigmoid 6 7 in the case of multilayer neural networks, Radial basis 8 etc). 9 These regression techniques can also be used to 10 11 construct a wavelet basis function set directly. 12 Methodologies for restricting the search space of the 13 wavelet basis functions considered are known. 14 this wavelet network has been shown to be effective for 15 chaotic time series prediction, its implementation 16 17 involves the use of wavelet frames of a decimated, dyadic, construction. The method of the present 18 invention may employ continuous wavelet networks 19 spanning a redundant wavelet basis which, although 20 computationally more expensive, overcomes the time 21 22 invariance constraint and the limited size of input 23space associated with use of wavelet frames. 24 25 The method may use conventional gradient decent methods 26 to produce a single layer wavelet classifier. 27 28 These wavelet networks may be further employed as part 29 of a multilayer system as a non-parameterised estimate of the original trace for input to further hidden 30 layers. 31

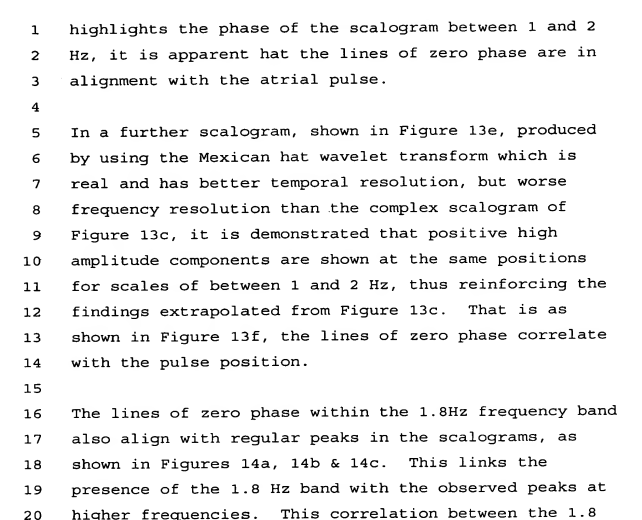


37 The network type of choice for the automated prediction 1 system of the method is selected on the basis of its 2 sensitivity and selectivity in correctly classifying 3 successful defibrillation outcomes in test set data, 4 since this is most clinically useful. 5 6 Thus experimental comparison of the three techniques 7 demonstrates the efficacy of the wavelet transform 8 9 technique. 10 The nature of underlying atrial activity can also be 11 determined from wavelet decomposition of the EKG 12 The wavelet function gives information 13 signal. regarding the amplitude and, where appropriate, phase 14 of the transformed signal. It is known that pressure 15 readings taken from the aorta correlate to forms of 16 atrial activity within the heart. Areas of localised 17 high energy contained within the scalogram can be 18 demonstrated to correlate with these pressure readings. 19 This experimental result is extrapolated to mean that 20 areas of localised high energy contained within the 21 22 scalogram correlate with forms of atrial activity within-the-heart. 2-3-24 Figure 13a shows the aorta pressure, Figure 13b the EKG 25 trace, for the same time period as Figure 13a, and 26 Figure 13c shows the scalogram for the EKG of Figure 27 It is apparent that there is an increase in 28 energy in the system during an atrial pulse, indicated 29 by the dark blotches occurring in the scalogram at an 30

 $f_{
m bpc}$ of around 10 Hz. There is a frequency component

between 1 and 2 Hz. As shown in Figure 13d, which

31



21 22

activity is present.

In a further application of the method, means for identifying the optimum timing for application of the defibrillation shock can be extrapolated from the pulsing identified by the wavelet technique and shown in Figures 8, 9, 10, and 14, by comparison with traces of attempts at defibrillation which initially fail but are subsequently successful.

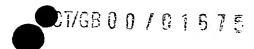
Hz band and the aorta pressure pulse suggests atrial





1 Thus, any data sets, in the above, that correspond to 2 3 multiple shocking of the same patient, where defibrillation has been repeatedly attempted are considered separately since these traces hold important 5 information. 6 8 The pilot study detailed above used Morlet wavelet based energy scalogram decomposition of signal segments 10 immediately prior to shocking. A full parametric 11 wavelet study of the method determines the optimum method. 12 13 The method includes the development of a classifier 14 using the wavelet transform analysis. 15 16 17 Various types of neural network classifier are achievable using this method. 18 19 20 The linkage of shock timing to the phase information of 21 wavelet components allows for increased defibrillation 22 success and reduced shock energies. The waveletderived information can also be employed to predict the 23 24 likelihood of shock success, preventing futile or harmful defibrillation attempts, and providing a 25 26 predictor of an optimal resuscitation strategy or 27 strategies. 28 29 This method demonstrates the utility of the wavelet 30 transform as a new method of EKG signal analysis during 31 It provides a robust, real-time solution to the





problem of useful monitoring of the myocardium during 1 2 resuscitation. When compared with conventional statistical methods, 4 such as fast Fourier transforms, it is seen that the 5 temporal resolution of the wavelet technique gives a 6 scalogram which better describes the non-stationary, 7 intermittent, nature of the EKG trace to be analysed, 8 and gives a method of greater predictive effectiveness 9 The effectiveness criteria for 10 than is already known. the networks of the method of the present invention are 11 12 based upon their sensitivity and selectivity in 13 correctly classifying successful defibrillation 14 outcomes from test data sets. 15 Although this description refers to wavelet transform 16 17 analysis, this term is to be construed to include 18 matching pursuit algorithms and similar analysis 19 techniques. 20 21 Modifications and improvements can be made to the above 22 without departing from the scope of the invention.

1	CLAI	MS
2		
3	1.	A method of decomposition of waveforms in a
4		cardiac signal using wavelet transform analysis.
5		
6	2.	A method as claimed in Claim 1 comprising the step
7		of employing discretized wavelet transform
8		analysis to process the said waveform.
9		
10	3.	A method as claimed in Claim 1 comprising the step
11		of employing discretized continuous wavelet
12		transform analysis to process the cardiac
13		waveform.
14		
15	4.	A method as claimed in any preceding claim
16		comprising the steps of deriving the wavelet
17		energy surfaces of an electrocardiogram (EKG)
18		signal; and plotting said wavelet energy surfaces
19		against a location parameter b, and a scale
20		parameter.
21		
22	5.	A method as claimed in Claim 4 wherein said scale
2:3		parameter is dilation a.
24		
25	6.	A method as claimed in Claim 4 wherein said scale
26		parameter is band pass frequency f_{bpc} .
27		
28	7.	A method as claimed in any preceding claim
29		comprising the initial steps of connecting
30		electrodes to a presenting patient; and sampling
31		the analogue input signals recorded to derive the
32		cardiac signal.



A method as claimed in any preceding claim 1 8. 2 including visually displaying the cardiac signal. 3 A method as claimed in any preceding claim 4 9. including visually displaying the distribution of 5 energies within the cardiac signal. 6 7 A method as claimed in any preceding claim 8 including visually displaying coherent structures 9 within the cardiac signal. 10 11 12 11. A method as claimed in any preceding claim including visually displaying the signal in real-13 14 time for clinical use. 15 A method as claimed in any preceding claim 16 12. comprising the step of constructing a contour plot 17 to display the decomposed waveform obtained. 18 19 A method as claimed in any preceding claim 20 13. comprising the step of constructing a surface plot 21 to display the decomposed waveform obtained. 22 23 24 14. A method as claimed in any preceding claim 25 comprising the step of constructing a 2D or a 3D energy scalogram to display the decomposed 26 27 waveform obtained.

28

15. A method as claimed in any preceding claim
 30 including the step of disassociating the component
 31 features of the temporal trace of a recorded EKG.



A method for the analysis of an EKG of a heart in 1 ventricular fibrillation including the method as 2 claimed in any preceding claim. 3 A method for the analysis of an EKG of a heart in 17. 5 ventricular fibrillation after the commencement of 6 7 cardio-pulmonary resuscitation (CPR) including the method as claimed in any of Claims 1 to 15. 9 A method as claimed in Claim 17 including the step 10 18. of temporal filtering of the EKG signal of a heart 11 that is subject to CPR to disassociate the CPR 12 signal from the heart signal. 13 14 A method as claimed in Claim 17 or Claim 18 using 15 19. 16 wavelet energy scalograms. 17 18 20. A method as claimed in Claim 17 or Claim 18 using 19 ridge following techniques 20 21 21. A method as claimed in Claim 20 wherein said ridge 22 following techniques are modulus maxima 23 techniques. 24 A method for the estimation of the health of a 25 22. heart in VF including the method of any of Claims 26 27 1 to 15 to provide measurable characteristics. 28 29 23. A method as claimed in Claim 22 wherein said

measurable characteristics are used to provide an estimate of the time elapsed since the onset of a cardiac incident.



•

A method as claimed in Claim 22 wherein said 24. 1 2 measurable characteristics are used to provide an 3 estimate of the health of a heart after commencement of CPR. 4 25. A method as claimed in any of Claims 22 to 24 6 7 wherein said measurable characteristics are used 8 to predict the outcome of a given therapeutic intervention. 9 10 11 26. A method as claimed in any of Claims 22 to 25 12 wherein said measurable characteristics are used to provide a guide for the optimal timing of 13 14 defibrillation of a heart in VF. 15 16 27. A method for the analysis of an EKG of a heart in 17 atrial fibrillation including the method as 18 claimed in any of Claims 1 to 14. 19 20 28. A method as claimed in Claim 27 including the step of partitioning the signal to provide separate 21 22 traces of QRS and T waves, and/or atrial activity 23 and/or background noise. 24 25 29. A method as claimed in any preceding claim 26 including the step of constructing a damage index 27 for reference purposes. 28 A method as claimed in Claim 29 wherein 29 30. 30 construction of said index includes the step of 31 developing network classifier from a library of recorded data. 32

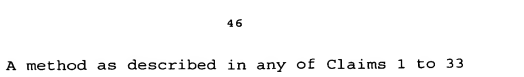




1	31.	A method as claimed in Claim 30 wherein said
2		network classifier developed is a neural network.
3		
4	32.	A method as claimed in any of Claims 29 to 31
5		wherein said network classifier developed is a
6		wavelet network classifier.
7		
8	33.	A method of decomposition of cardiac waveforms
9		using matching pursuit algorithms.
10		
11	34.	Apparatus for decomposition of waveforms in a
12		cardiac signal, said apparatus comprising wavelet
13		transform analysis means.
14		
15	35.	Apparatus as claimed in Claim 34 including means
16		to display the distribution of energies within a
17		waveform.
18		
19	36.	Apparatus as claimed in Claim 34 or Claim 35
20		including a monitor adapted to display decomposed
21		waveforms.
22		·
2:3	37.	Apparatus as claimed any of Claims 34 to 36
24		adapted for inclusion in an EKG apparatus.
25		
26	38.	Defibrillation means adapted to operate in
27		response to a signal generated by comparison of an
28		EKG trace with decomposed waveform obtained by the
29		method of any of Claims 1 to 33.



with reference to or as shown in the accompanying



39.

drawings.

Apparatus as described in any of Claims 34 to 38 40. with reference to or as shown in the accompanying drawings.

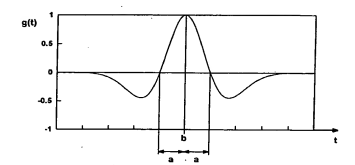
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3 73 73 75	RACT
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- 3 A method of analysis of medical signals which uses
- 4 wavelet transform analysis to decompose cardiac
- 5 signals. Apparatus for carrying out the method, and
- 6 cardiac apparatus adapted to employ the method are also
- 7 described.



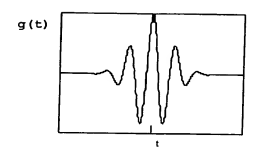


Figure 1(a)

Figure 1(b)

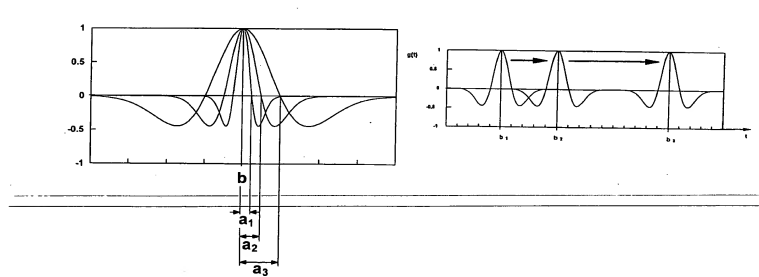


Figure 2(a)

Figure 2(b)

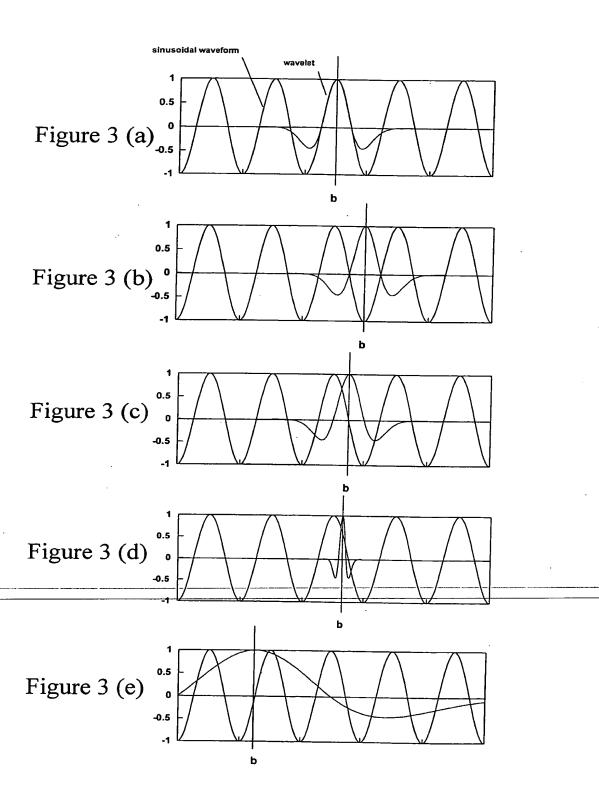


Figure 4 (a)

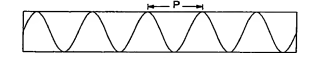


Figure 5 (a)

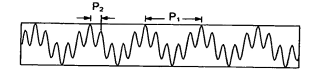


Figure 4 (b)

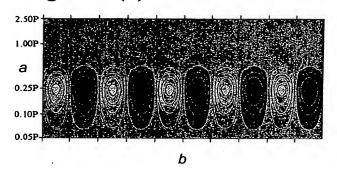


Figure 5 (b)

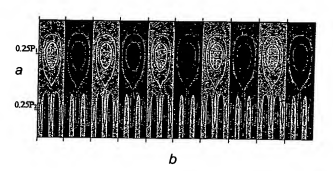


Figure 4 (c)

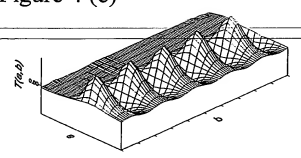


Figure 5 (c)

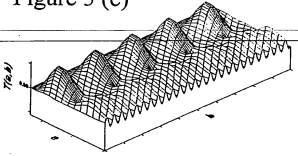


Figure 6 (a)

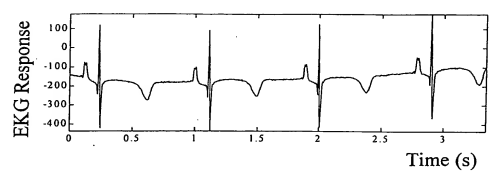


Figure 6 (b)

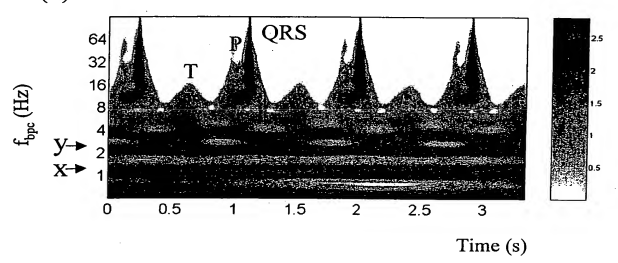
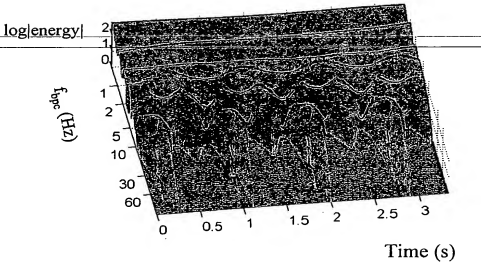
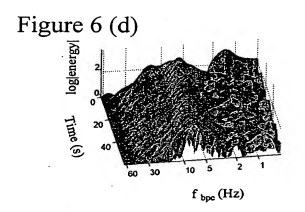
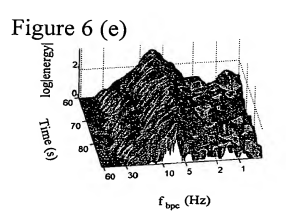
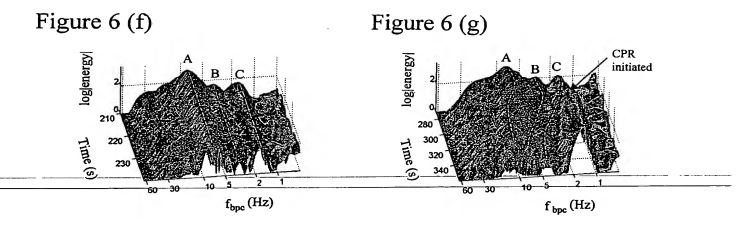


Figure 6 (c)









Time (minutes)

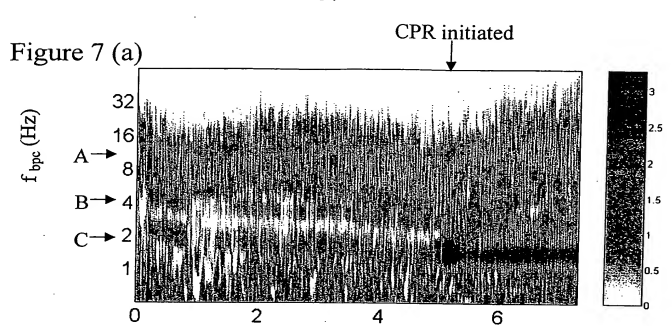


Figure 7 (b)

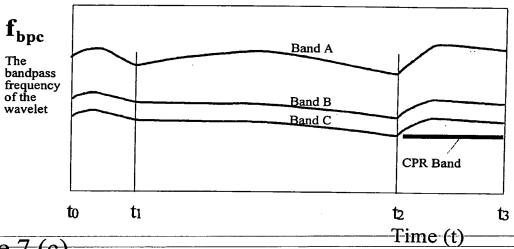
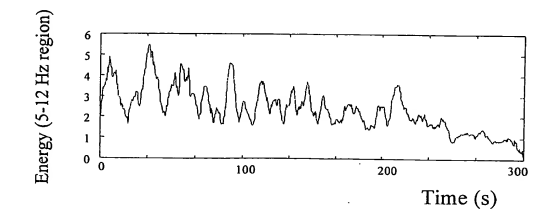
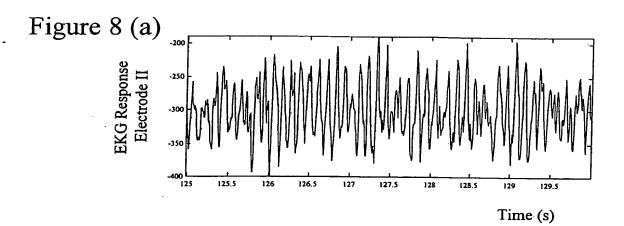
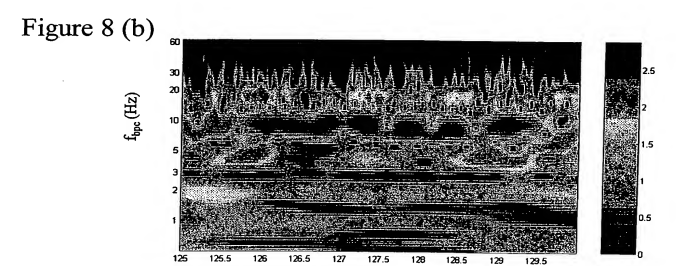


Figure 7 (c)

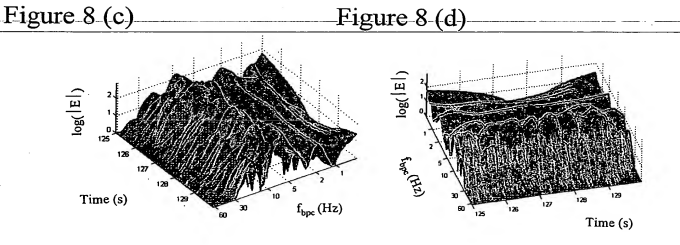




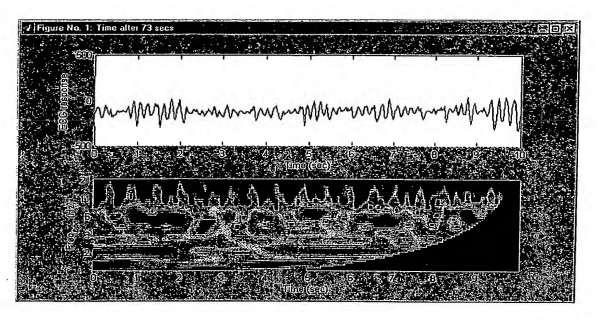


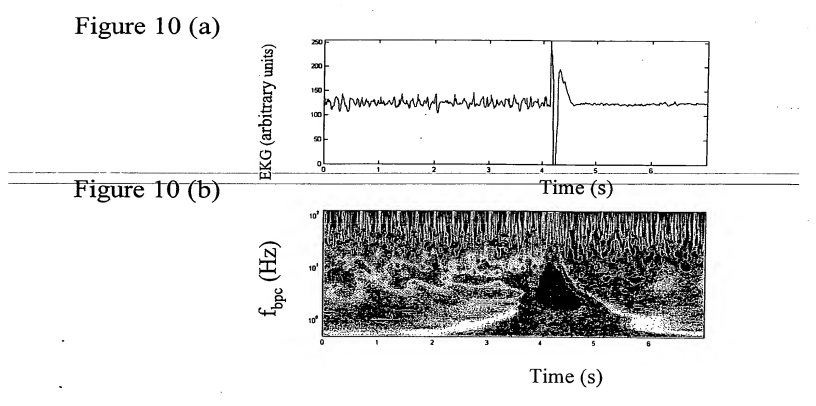


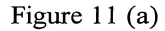
Time (s)











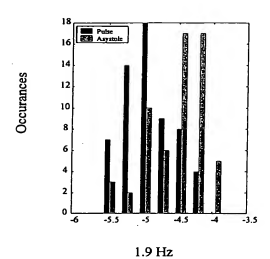
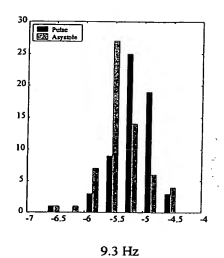
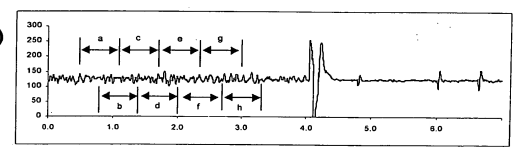


Figure 11 (b)



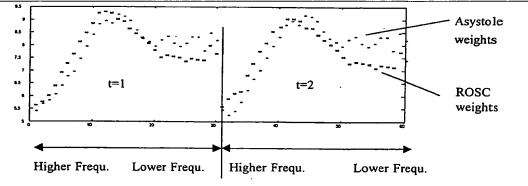
mean energy (log)

Figure 12 (a)

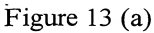


Weight value

Figure 12 (b)







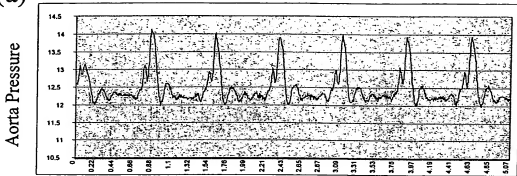


Figure 13 (b)

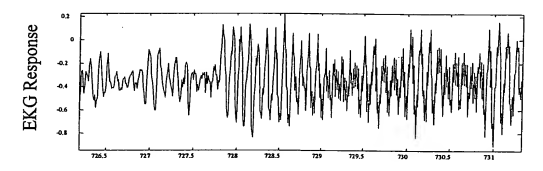


Figure 13 (c)

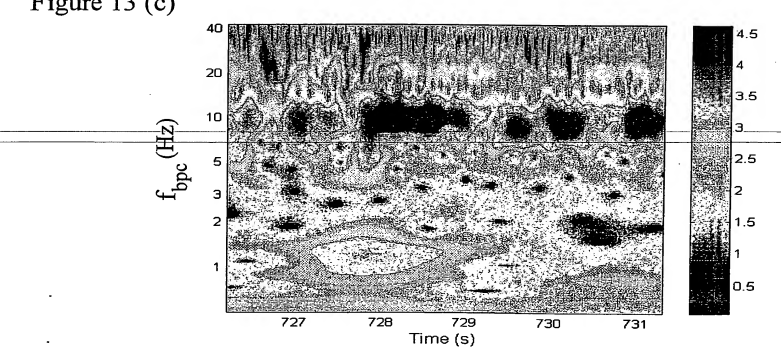




Figure 13 (d)

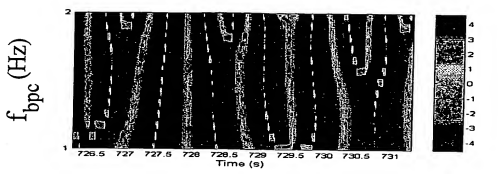


Figure 13 (e)

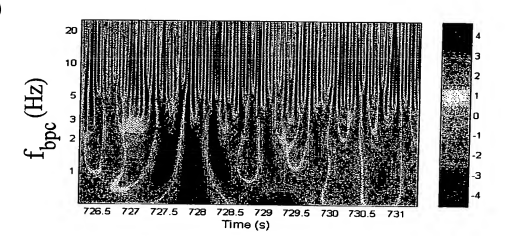
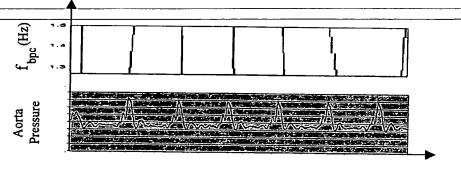
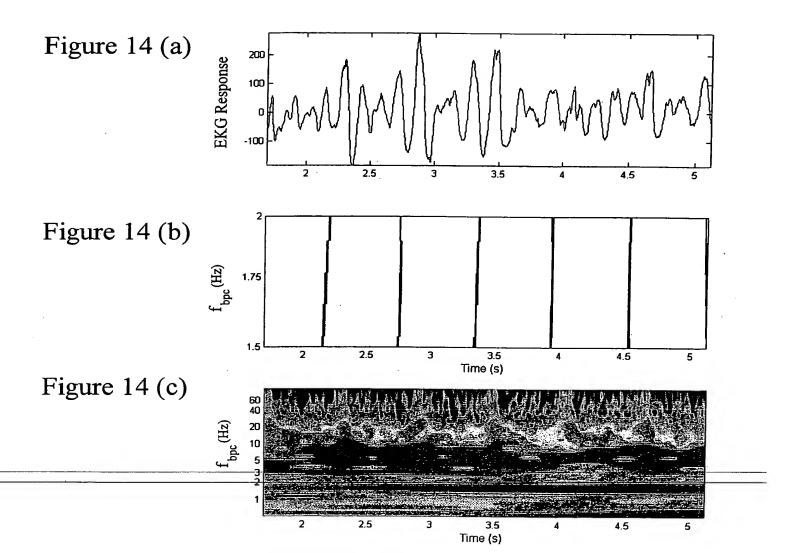


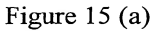
Figure 13 (f)

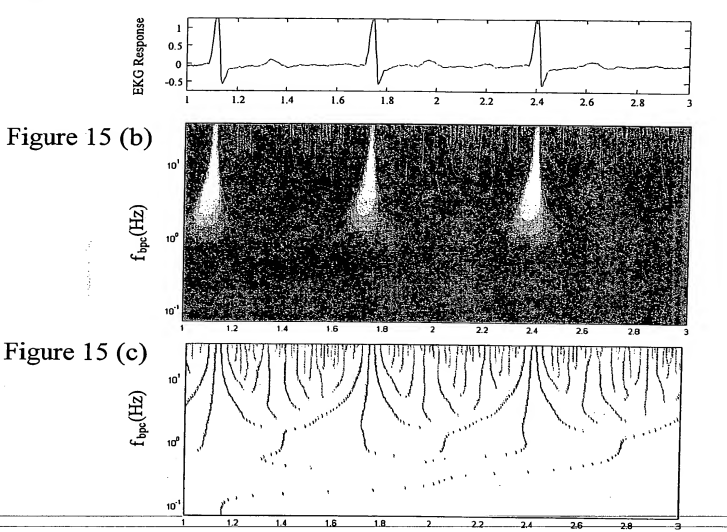


Time (s)

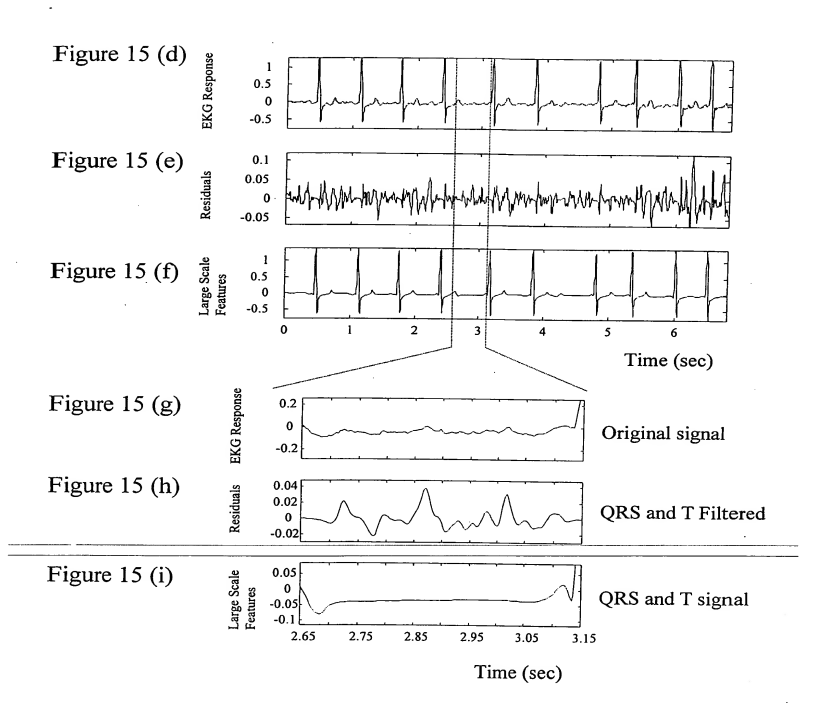








Time (Seconds)



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REC'D 22	DEC 2000
WIPO	PCT

INTERNATIONAL SEARCH REPORT

(PCT Article 18 and Rules 43 and 44)

Applicant's or agent's file reference P23847A/JMK	FOR FURTHER see Notification (Form PCT/ISA/	of Transmittal of International Search Report 220) as well as, where applicable, item 5 below.
International application No.	International filing date (day/month/year)	(Earliest) Priority Date (day/month/year)
ппенавона аррисавон но.	The mail of the fact (day, more you,	
PCT/GB 00/01675	02/05/2000	01/05/1999
Applicant		
THE COURT OF NAPIER UNIV	ERSITY.	
This International Search Report has be according to Article 18_A copy is being	en prepared by this International Searching Autransmitted to the International Bureau.	thority and is transmitted to the applicant
·		
This International Search Report consis		- roped
X It is also accompanied t	by a copy of each prior art document cited in this	s report.
Basis of the report		
•	e international search was carried out on the ba	sis of the international application in the
language in which it was filed, u	nless otherwise indicated under this item.	.,
the international search	was carried out on the basis of a translation of	the international application furnished to this
Authority (Rule 23.1(b))		
b. With regard to any nucleotide a was carried out on the basis of t	and/or amino acid sequence disclosed in the i	nternational application, the international search
	tional application in written form.	
filed together with the in	ternational application in computer readable for	m.
furnished subsequently	to this Authority in written form.	
furnished subsequently	to this Authority in computer readble form.	
the statement that the s	ubsequently furnished written sequence listing of as filed has been furnished.	does not go beyond the disclosure in the
the statement that the in furnished	nformation recorded in computer readable form	is identical to the written sequence listing has been
2. X Certain claims were fo	ound unsearchable (See Box I).	
3. X Unity of invention is la	cking (see Box II).	
4. With regard to the title,		
X the text is approved as	submitted by the applicant.	
the text has been estab	lished by this Authority to read as follows:	
5. With regard to the abstract,		
	submitted by the applicant.	
the text has been estab within one month from t	lished, according to Rule 38.2(b), by this Author he date of mailing of this international search re	ity as it appears in Box III. The applicant may, port, submit comments to this Authority.
	blished with the abstract is Figure No.	9
X as suggested by the ap		None of the figures.
	ailed to suggest a figure.	
	er characterizes the invention.	
because this lightle bett	or characterizes the invention.	



Intel Ponal application No. PCT/GB 00/01675

Box I Observations where certain claims were found unsearchable (Continuation of item 1 of first sheet)
This International Search Report has not been established in respect of certain claims under Article 17(2)(a) for the following reasons:
Claims Nos.: because they relate to subject matter not required to be searched by this Authority, namely:
2. X Claims Nos.: Because they relate to parts of the International Application that do not comply with the prescribed requirements to such an extent that no meaningful International Search can be carried out, specifically: See FURTHER INFORMATION sheet PCT/ISA/210
3. Claims Nos.: because they are dependent claims and are not drafted in accordance with the second and third sentences of Rule 6.4(a).
Box II Observations where unity of invention is lacking (Continuation of item 2 of first sheet)
This International Searching Authority found multiple inventions in this international application, as follows:
see additional sheet
As all required additional search fees were timely paid by the applicant, this International Search Report covers all searchable claims.
2. As all searchable claims could be searched without effort justifying an additional fee, this Authority did not invite payment of any additional fee.
3. As only some of the required additional search fees were timely paid by the applicant, this International Search Report covers only those claims for which fees were paid, specifically claims Nos.:
—— covers only those claims for which lees were paid, specifically claims Nos
4. No required additional search fees were timely paid by the applicant. Consequently, this International Search Report is restricted to the invention first mentioned in the claims; it is covered by claims Nos.: 1-16, 33-38
Remark on Protest The additional search fees were accompanied by the applicant's protest. No protest accompanied the payment of additional search fees.

FURTHER INFORMATION CONTINUED FROM PCT/ISA/ 210

This International Searching Authority found multiple (groups of) inventions in this international application, as follows:

1. Claims: 1-16, 33, 34-38

Method and device for ECG analysis using wavelet transformation or matching pursuit algorithms and visually displaying the signal and/or the decomposed waveform.

2. Claims: 17-21

Analysis of an ECG of a heart in ventricular fibrillation after commencement of CPR and method of disassociating the CPR signal from the heart signal.

3. Claims: 22-32

Method of estimating the health of a heart in ventricular fibrillation in order to guide therapeutic intervention or to predict outcome.

FURTHER INFORMATION CONTINUED FROM PCT/ISA/ 210

Continuation of Box I.2

Claims Nos.: 39 40

Rule 6.2 (a)

References to Other Parts of the International Application Claims shall not, except where absolutely necessary, rely, in respect of the technical features of the

invention, on references to the description or drawings. In particular, they shall not rely on such

references as: "as described in part ... of the description," or "as illustrated in figure ... of the drawings."

The applicant's attention is drawn to the fact that claims, or parts of claims, relating to inventions in respect of which no international search report has been established need not be the subject of an international preliminary examination (Rule 66.1(e) PCT). The applicant is advised that the EPO policy when acting as an International Preliminary Examining Authority is normally not to carry out a preliminary examination on matter which has not been searched. This is the case irrespective of whether or not the claims are amended following receipt of the search report or during any Chapter II procedure.

A. CLASSIFICATION OF SUBJECT MATTER IPC 7 G06F17/00

According to International Patent Classification (IPC) or to both national classification and IPC

B. FIELDS SEARCHED

Minimum documentation searched (classification system followed by classification symbols) I PC $\,7\,$ G06F $\,$ A61N

Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched

Electronic data base consulted during the international search (name of data base and, where practical, search terms used)

EPO-Internal, WPI Data, PAJ, INSPEC

	On the selection of the relevant page 2000	Relevant to claim No.
Category °	Citation of document, with indication, where appropriate, of the relevant passages	Tiolovani io diamini
X	WO 96 08992 A (SHOSHAN HERBERT Z ;UNIV	1,2,7-9,
•	RAMOT (IL); AKŠELROD SOLANGE (IL);	12-15, 34-37
٨	KESELBR) 28 March 1996 (1996-03-28) page 9, line 27 -page 11, line 9	3-5,10,
Α	page 9, Title 27 -page 11, Title 9	11,16
	page 13, line 29 -page 14, line 14	
X	US 5 439 483 A (DUONG-VAN MINH)	1,2,7,
	8 August 1995 (1995-08-08)	16,22,34 3,23,
Α	column 4, line 36-49	25-27,38
	-/	
	•	

X Further documents are listed in the continuation of box C.	X Patent family members are listed in annex.
"A" document defining the general state of the art which is not considered to be of particular relevance "E" earlier document but published on or after the international filing date "L" document which may throw doubts on priority claim(s) or which is cited to establish the publication date of another citation or other special reason (as specified) "O" document referring to an oral disclosure, use, exhibition or other means "P" document published prior to the international filing date but later than the priority date claimed	"T" later document published after the international filing date or priority date and not in conflict with the application but cited to understand the principle or theory underlying the invention "X" document of particular relevance; the claimed invention cannot be considered novel or cannot be considered to involve an inventive step when the document is taken alone "Y" document of particular relevance; the claimed invention cannot be considered to involve an inventive step when the document is combined with one or more other such documents, such combination being obvious to a person skilled in the art. "&" document member of the same patent family
Date of the actual completion of the international search	Date of mailing of the international search report
13 September 2000	2 2. 12. 2000
Name and mailing address of the ISA	Authorized officer
European Patent Office, P.B. 5818 Patentlaan 2 NL - 2280 HV Rijswijk Tel. (+31-70) 340-2040, Tx. 31 651 epo nl, Fax: (+31-70) 340-3016	Gro∌mann, C

4



	DOCUMENTS CONSIDERED TO BE RELEVANT tion of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
Category ° Cita	non or document, with indication, where appropriate, or the relevant passages	Televant to Gain 140.
	DATABASE INSPEC [Online] INSTITUTE OF ELECTRICAL ENGINEERS, STEVENAGE, GB; 13 September 1998 (1998-09-13) MILLET-ROIG J; LOPEZ-SORIANO JJ; MOCHOLF A ET AL.: "Study of frequency and time domain parameters extracted by means of wavelet transform applied to ECG to distinguish between VF and other arrhythmias" XP002145546 abstract	1,2,16
	CHEN J ET AL: "ECG DATA COMPRESSION BY USING WAVELET TRANSFORM" IEICE TRANSACTIONS ON INFORMATION AND SYSTEMS, JP, INSTITUTE OF ELECTRONICS INFORMATION AND COMM. ENG. TOKYO, vol. E76-D, no. 12, 1 December 1993 (1993-12-01), pages 1454-1461, XP000435570 ISSN: 0916-8532 abstract	1,2
	DATABASE INSPEC [Online] INSTITUTE OF ELECTRICAL ENGINEERS, STEVENAGE, GB; 5 November 1996 (1996-11-05) GEVA A B: "Spatio-temporal matching pursuit (SToMP) for multiple source estimation of evoked potentials" XP002145547 abstract	33
	SAVA H ET AL: "APPLICATION OF THE MATCHING PURSUIT METHOD FOR STRUCTURAL DECOMPOSITION AND AVERAGING OF PHONOCARDIOGRAPHIC SIGNALS" MEDICAL AND BIOLOGICAL ENGINEERING AND COMPUTING,GB,PETER PEREGRINUS LTD. STEVENAGE, vol. 36, no. 3, 1 May 1998 (1998-05-01), pages 302-308, XP000751653	33
A	ISSN: 0140-0118 the whole document US 5 795 304 A (LEE KAE YOL ET AL) 18 August 1998 (1998-08-18) column 6, line 12 -column 7, line 23	1,3

4



pplication No

PCT/GB 00/01675

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WO 9608992	Α	28-03-1996	AU 3717495 A EP 0869734 A JP 11511036 T US 5797840 A	09-04-1996 14-10-1998 28-09-1999 25-08-1998
US 5439483	Α	08-08-1995	NONE	
US 5795304	Α	18-08-1998	NONE	